

THE CONTROL OF GROWTH & DIFFERENTIATION IN PLANTS

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**THE CONTROL OF GROWTH AND
DIFFERENTIATION IN PLANTS**

SECOND EDITION



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THE CONTROL OF
GROWTH AND
DIFFERENTIATION
IN PLANTS

SECOND EDITION

by

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To Helen and Anne

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Preface to the Second Edition

THE STEADY increase in our knowledge and understanding of plant development which has occurred since the first edition of this book almost 10 years ago has necessitated its extensive revision. In the present edition large sections have been rewritten and extended to bring the information up to date, and an additional chapter on Phytochrome and Photomorphogenesis is now included. There has also been extensive rearrangement of the material to give a more logical and coherent presentation.

We are indebted to our colleagues Dr. M. A. Hall and Dr. P. F. Saunders, and to Professor J. Heslop-Harrison and Professor J. Zeevaart, for reading various sections of the revised manuscript and for their helpful and constructive criticisms.

Preface to the First Edition

THE PHENOMENON of development, in both plants and animals, presents some of the most challenging unsolved problems of biology and is one of the remaining areas in which it cannot yet be said that we have made a decisive "breakthrough", comparable with the recent advances in biochemistry and molecular biology. Nevertheless, during the past 30 or 40 years there has been a steady advance in our understanding of the physiology of growth and differentiation in higher plants, so that there is now a considerable body of well-established knowledge in this field. It is generally accepted that some knowledge of the physiology of plant growth and differentiation is important for all students of the botanical sciences, and sections on this subject are commonly included in textbooks of plant physiology. The developmental approach is now also given more prominence in the contemporary teachings of plant form and structure than in the past, but there have been few attempts to bring together both morphological and physiological approaches within the same volume, and this we have attempted to do. Unless we attempt to relate the two approaches to each other, morphological and anatomical accounts of growth and differentiation must remain largely descriptive in nature, whereas our aim should clearly be to understand the processes underlying and controlling the structural changes. Conversely, physiological and biochemical studies which are not related back to developmental processes in the plant are liable to lose relevance and biological significance.

While it is true that we have found it convenient to concentrate on the structural aspects of development in the first few chapters, nevertheless we have attempted, throughout the book, to base our whole approach upon the growth and differentiation of organs and tissues, and the major developmental changes in the plant as a whole. Thus, after giving a brief account of the chemistry and biochemistry of plant hormones, we return to consider their role in the control of growth and differentiation at various levels. We then consider the factors regulating the major phase-switches in the development of the plant as a whole, viz. those controlling flowering, dormancy and senescence.

Finally, in the last chapter, we examine some basic problems in plant development, including the nature of the control mechanisms in differentiation. There are large gaps in our understanding of some of the most important aspects of development, such as the nature of the processes controlling the orderly sequence of changes so characteristic of development of all organisms. The discussion on such topics is, of necessity, largely speculative, but current thought in molecular genetics and theories of gene activation and repression derived

from studies on micro-organisms enable us to formulate the problems in more precise terms than has been possible hitherto.

The book is intended primarily as an introduction to plant growth and differentiation for undergraduate students. While we have attempted, so far as possible, to give the supporting evidence for statements, references to the individual pieces of research upon which we have drawn are reduced to a minimum. We believe that an over-extensive chronicle of research workers' names and dates only serves to distract the student's attention from the main theme. We are confident that the many unnamed researchers whose work has made this book possible will not be offended. To assist the student, however, we have provided a list of additional reading, from which he can obtain detailed reference lists.

This book has only been made possible by the kind assistance of many persons in different ways and we make grateful acknowledgement of their help. We should mention specially, however, Professor A. W. Galston, Professor H. Heslop-Harrison, our editors, Professor G. F. Asprey and Dr. A. G. Lyon, and our colleagues, Dr. M. A. Hall and Dr. P. F. Saunders, all of whom have read sections of the book and have made most helpful suggestions. Responsibility for any errors which may have crept into the book must remain ours, however. We also wish to thank Miss M. Bigwood for her skilled assistance in the preparation of some of the diagrams.

CHAPTER I

Growth in the Higher Plant

INTRODUCTION

We are all so familiar with the remarkable changes which occur during the life cycle of a plant, from germination to fruiting and senescence, that we tend to take the phenomenon of development for granted, so that it may cease to excite our wonder. Nevertheless, the orderly succession of changes leading from the simple structure of the embryo to the highly complex organization of the mature plant presents some of the most fascinating and challenging outstanding problems of biology. In this book we shall be concerned primarily with describing and examining what is known about the processes underlying and controlling plant development.

Before we can proceed, however, it is necessary to define certain terms, which are not always used in precise senses. *Development* is applied in its broadest sense to the whole series of changes which an organism goes through during its life cycle, but it may equally be applied to individual organs, to tissues or even to cells. Development is most clearly manifest in changes in the form of an organism, as when it changes from a vegetative to a flowering condition. Similarly, we may speak of the development of a leaf, from a simple primordium to a complex, mature organ.

Plant development involves both *growth* and *differentiation*. The term growth is applied to *quantitative* changes occurring during development and it may be defined as an irreversible change in the size of a cell, organ or whole organism. The external form of an organ is primarily the result of *differential growth* along certain axes. However, during development there appear not only quantitative differences in the numbers and arrangement of cells within different organs, but also *qualitative* differences between cells, tissues and organs, to which the term *differentiation* is applied. Differentiation at the cell and tissue level is well known and is the primary object of study in plant anatomy. However, we may also speak of differentiation of the plant body into shoot and root. Similarly, the change from the vegetative to the reproductive phase may also be regarded as another example of differentiation. We shall, therefore, apply the term differentiation in a very broad sense to any situation in which meristematic cells give rise to two or more types of cell, tissue or organ which are qualitatively different from each other.

Thus, we may say that *growth and differentiation are the two major developmental processes*. Usually growth and differentiation take place concurrently during development, but under certain conditions we may obtain growth without differentiation, as in the growth of a mass of callus cells (Chapter 6).

The problems of development can be studied in a number of different ways, but basically there are two major types of approach, viz. (1) the morphological and (2) the physiological and biochemical. Developmental morphology and anatomy were formerly largely concerned with describing the visible changes occurring during development, but current interest is mainly directed to trying to understand the factors and processes determining plant form, using experimental techniques, such as surgery, tissue culture, autoradiography and so on. However, development cannot be fully understood without a study of the manifold biochemical and physiological processes underlying and determining the morphological changes, and it is these latter aspects of development which form the main subject of this work.

The experimental morphologist often uses the term *morphogenesis*, which, in the literal sense, is concerned with the origin of form in living organisms. However, by the term "form" should be understood not only the gross external morphology of the plant, but its whole organization, which may be recognized as existing at several different levels; thus, we may recognize (1) the structural organization of the individual cell, as shown by electron microscopy, (2) the organization of cells to form tissues, and (3) the organization of the plant body at the macroscopic level. Moreover, in the study of morphogenesis we are concerned not only with observable changes in form and structure but also with the underlying processes controlling the development of organs and tissues, and insofar as these processes must ultimately be explainable in terms of physics and chemistry, this aspect of morphogenesis is identical with developmental physiology and biochemistry. However, at the present time our knowledge of the molecular basis of morphogenesis is very fragmentary and we know very little about the physiological and biochemical processes regulating, for example, the initiation and development of leaves.

When we come to consider the physiology of development, we find a further dichotomy of approach. On the one hand, a considerable body of knowledge has been acquired about the role of hormones as "internal" factors controlling growth and differentiation; on the other hand, the profound importance of environmental factors, such as day length and temperature, in the regulation of some of the major phases in the plant life cycle has been clearly demonstrated, although there is considerable evidence that a number of environmental influences are mediated through effects on the levels and distribution of hormones within the plant.

It is axiomatic that the plant body at any given stage is the resultant of the interaction between the inherent (genetic) potentialities of the species and the external factors of the environment. Thus, we cannot say that certain characteristics of the plant are determined genetically, whereas others are environmentally determined, since *all* its characteristics are affected by both genetical and environmental influences. However, it is quite legitimate to say that some *differences* between plants are primarily genetically determined whereas

others are due to environmental factors. Thus, the lack of chlorophyll in a plant may be caused by a mutation affecting chlorophyll biosynthesis. On the other hand, a plant may lack chlorophyll because it has been grown in the dark, so that it is etiolated. But it must be emphasized again that the development of a normal green leaf requires both the appropriate genetical factors and certain environmental conditions, including light.

When we speak of the genetical potentialities of the species we must include not only genes located in the nucleus, but also cytoplasmic factors. Certain characters of the plant, including some chloroplast characters, show cytoplasmic inheritance. This fact should not surprise us unduly, since it is now well established that chloroplasts contain DNA and are probably self-replicating organelles. In this book we are not primarily concerned with genetical aspects of morphogenesis, but in all our discussions of this problem it is a basic assumption that, in the final analysis, development involves the expression of the information stored in the genes.

THE LOCALIZATION OF GROWTH

One of the essential characteristics of organisms is that they are able to take up relatively simple substances from their environment and use them in the synthesis of the varied and complex substances of which cells are composed. It is this increase in the amount of living material which is basically what we mean by growth. At the cellular level the increase in living material normally leads to an increase in cell size and ultimately to cell division. These two aspects of growth are seen in their simplest form in unicellular organisms such as bacteria, unicellular algae and protozoa, where growth leads to enlargement of each cell which then divides and the process is repeated.

When we come to consider the growth of multicellular organisms, such as the higher plants, the situation is much more complex. It is true that here, also, growth ultimately depends on the enlargement and division of individual cells, but not all cells of the plant body contribute to the growth of the organisms as a whole, for growth is restricted to certain embryonic regions, the *meristems*. This restriction of the growing regions is probably related to the fact that mature plant cells are normally surrounded by relatively thick and rigid cell walls, and many cells of mechanical and vascular tissues are, of course, non-living. These facts would probably render co-ordinated growth, involving both cell division and cell enlargement, difficult in an organ, such as a stem once a certain stage of differentiation had been reached. We shall see later that most living plant cells retain the capacity to divide under certain conditions, but even if they do divide the daughter cells do not necessarily increase in size, unless they are relatively thin-walled cells which are able to revert to the embryonic or "meristematic" condition. In having rather strictly localized embryonic regions higher plants differ from animals, where growth typically occurs throughout the organism as a whole.

This difference between higher plants and animals is no doubt related to the basic differences in the modes of nutrition of the two groups. Because they have to take up water and

mineral salts from the soil, the autotrophic land plants must necessarily be rooted and sessile, whereas most animals have to forage for their food, whether they are herbivorous or carnivorous, and they need, therefore, to be mobile. This requirement for mobility in animals which forage for their food, in turn, demands that they should have flexible bodies, whereas the plant body can be much more rigid and indeed it needs to be so in erect-growing plants, especially in large forest trees. This rigidity and firmness of the plant body depends upon the presence of relatively thick and firm cell walls, whether in the living cells of the leaf, for example, or in the non-living cells of mechanical tissue of the stem. (The rigidity of those tissues consisting mainly of living cells depends, of course, on the turgidity of the cells and not simply on the mechanical properties of the walls, but even in such tissues a cell wall is an essential requirement for the attainment of the turgid condition.) On the other hand, in aquatic plants, whether they are lower plants or angiosperms, nutrients may be absorbed from the surrounding water directly into the shoot, so that they may be free-floating, and the mechanical tissues are usually less well developed than in land plants.

A number of different types of meristem may be recognized in the plant body. The axial organs, the stems and roots, have *apical meristems*, i.e. growth in length is restricted to the tip regions and the new tissue is added to the plant body on the proximal side, so that the pattern of growth may be described as *accretionary*. The apical meristems of the stem and root usually remain permanently embryonic and capable of growth over long periods—for hundreds of years in some trees. Consequently we may describe these as *indeterminate meristems*.

On the other hand, other parts of the plant, particularly the leaves, flowers and fruits, show rather different patterns of growth and they are embryonic for only a limited period before the whole organ attains maturity. Thus, the growing regions of such organs are sometimes referred to as *determinate meristems*. In such organs the pattern of growth resembles that of animals in that, firstly, there is an embryonic phase of limited duration and secondly, in such organs growth is more generalized than in stems and roots.

The presence of indeterminate meristems, together with the capacity for forming branches, each with its apical meristem, gives the plant body a much less precise and definite form than is the case for the animal body. Indeed, the general form of the plant body resembles a colony of coelenterates, such as corals, rather than that of an individual higher animal. On the other hand, the organs showing determinate growth, such as leaves and flowers, generally show much more precise morphology and may have fairly precise numbers of parts, such as petals.

In addition to classifying meristems as indeterminate and determinate we may classify them in various other ways. For example, we may distinguish the apical meristems of stems and roots, from the *lateral meristems*, comprising the cambium and phellogen (cork cambium). In some plants there are *intercalary meristems*, inserted between regions of differentiated tissues. One of the best known examples of this type of meristem is seen in grasses, where the internodes and leaf sheaths continue growth in the basal region, after the upper parts have become differentiated. The structure of some of these meristems will be described in more detail in Chapter 2.

CELL DIVISION AND CELL VACUOLATION

The growth of a multicellular plant involves both increase in cell number, by cell division, and increase in cell size. These two aspects of growth have no sharp spatial boundaries; however, in the apical regions of shoots and roots cell division occurs most intensively towards the extreme tip of both organs, whereas the region of most rapid increase in cell size is in a zone a few millimetres back from the tip (Fig. 1.1). In organs of determinate growth, such as leaves and fruits, these two aspects of growth tend to be separated in time, so that there is an early phase in which cell division is predominant, followed later by a phase when cell division ceases and there is active increase in cell size. The greater part of this increase in size is due to vacuolation, i.e. by water uptake, and as a result the cytoplasm may come to be limited to a thin boundary layer against the cell wall.

In the tip regions of roots and shoots in which cell division predominates, the cells are relatively small, and have prominent, spherical nuclei lying towards the centre of the cytoplasm, which is non-vacuolated and tends to be densely staining; the cell walls are thin. The details of the process of mitosis by which the nucleus divides need not be described here. As a result of division, each of the two daughter cells is only half the size of the parent cell. These cells then proceed to enlarge, but such cell growth involves the synthesis of cytoplasm and cell wall material and not vacuolation.

Since the number of cells in the zone of cell division tends to remain fairly constant (at least over limited periods), it is clear that not all the daughter cells formed in this zone retain the capacity for unlimited further division. The situation is perhaps best illustrated by reference to plants which grow by a single apical cell, such as certain algae and the bryophytes and some pteridophytes where it can clearly be seen that division of the apical cell results in one cell on the outside which becomes the new apical cell, and a second daughter cell, on the proximal side, which gives rise to the differentiated tissue of the thallus or shoot. This latter daughter cell usually undergoes several further divisions but ultimately the derivative cells lose their capacity for division. Thus, whereas the apical cell remains permanently meristematic, the derivative cells are capable of only a limited number of further divisions. The situation must be analogous in the more complex apices of gymnosperms and angiosperms, where there is normally a number of initial cells, i.e. cells which remain meristematic and undergo repeated division, but it is more difficult to recognize which of the daughter cells is destined to remain meristematic and which will give rise to mature tissue. The problem of why cells in the initial zone remain permanently embryonic or meristematic, whereas the derivative cells on the proximal side are capable of only a limited number of further divisions is an intriguing one, but it remains unsolved at the present time.

At a certain distance from the apex, in both shoots and roots, the process of vacuolation commences and, as a result of this process, the root cells of *Allium* may increase in length from 17 μm to 30 μm and in volume by 30-fold. In other tissues the cells may increase up to 150-fold in total volume during vacuolation. It appears that this great uptake of water during cell extension is essentially governed by osmosis, and if we apply the usual concept relating to water uptake by cells, then, in general, the ability of the cell to take up water is

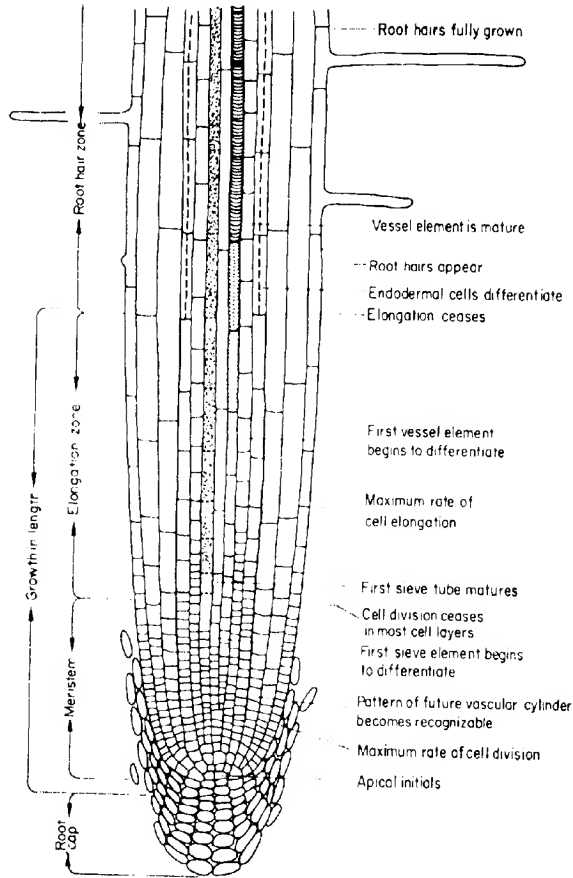


FIG. 1.1. Simplified diagram of the growing zone of a root, in longitudinal section. The number of cells in a living root is normally much greater than is shown in this diagram. (Reprinted from P. M. Ray, *The Living Plant*, Holt, Rinehart & Winston, New York and London, 1963.)

given by its water potential (ψ) which is equal to the osmotic potential (π) of the vacuolar solution plus the wall or turgor pressure (p). That is, $\psi = p + \pi$. Now clearly water uptake may involve changes in either the osmotic potential or in the wall pressure, or both. Studies on the changes in osmotic potential of the vacuolar solution during growth have yielded no evidence of a change in osmotic potential. Indeed, since the vacuolar sap becomes greatly diluted during growth, considerable amounts of additional osmotically active substances, such as sugars, salts, organic acids, etc., must pass into the vacuole during growth, in order simply to maintain the osmotic potential at a steady value. In some organs the osmotic potential of the vacuole may actually rise during this phase of growth. Thus, in the petioles of the water lily, *Victoria regia*, which may increase in length from 9 cm to 68 cm in 24 hours, the osmotic value may rise to less than half its original value during the extension phase. On the other hand, there is considerable evidence that in vacuolating cells the wall pressure is reduced by increased plasticity of the cell wall at this time (p. 83). As a result of its increased plastic extensibility the wall undergoes irreversible elongation during vacuolation.

Although the greater part of the increase in cell volume during vacuolation is due to water uptake, the synthesis of new cytoplasm and cell wall material proceeds actively during this period, so that the cell increases considerably in dry weight (Fig. 1.2). Thus, the

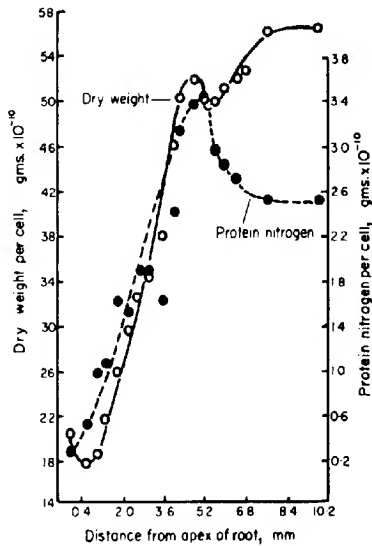


FIG. 1.2. Changes in dry weight and protein content of cells at increasing distances from the apex of pea roots. (Adapted from R. Brown and D. Broadbent, *J. Exp. Bot.* 1, 249-63, 1950.)

processes of cell growth initiated before vacuolation commences are continued during this latter phase. Moreover, the zones of cell division and cell vacuolation are not sharply demarcated, and in both shoots and roots of many species cell division occurs in cells which have started to undergo considerable vacuolation (Fig. 1.3). Division may also occur in vacuolated cells in wound tissues. In root tips the separation of the zones of division and vacuolation are somewhat sharper and division in vacuolated cells is less frequent.

Since growth involves various endergonic, i.e. energy-requiring processes, including protein synthesis, it is not surprising to find that rapidly elongating tissues of the root have a high respiration rate, when compared with mature tissues on the basis of equal volumes of tissue, although when expressed *per cell* the respiration rate of mature cells may be greater than that of meristematic cells, since the latter are smaller and contain less cytoplasm. Moreover, growth requires aerobic conditions and an adequate supply of carbohydrate, both as an energy source and as structural material.

The role of growth hormones in cell division and cell extension will be discussed later.

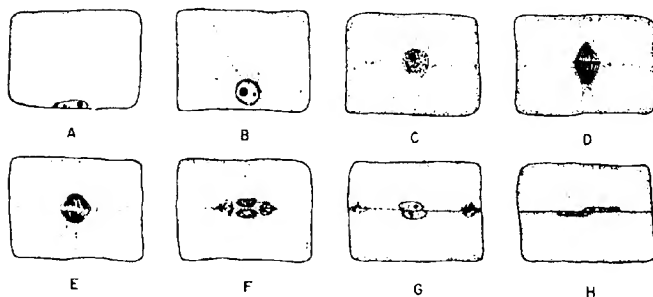


FIG. 1.3. Cell division in vacuolated cells. A, interphase; B, early prophase; C, prophase; D, metaphase; E, anaphase; F and G, telophases; H, two daughter cells at interphase. (From E. W. Sinnott and R. Bloch, *Amer. J. Bot.* 28, 1941.)

GROWTH OF CELL WALLS

During cell extension the area of the cell wall may increase greatly and this fact poses a number of problems. It might be expected that as the wall is stretched by turgor pressure, it would decrease in thickness, but usually this does not occur. Hence, new material must be added to the wall during growth. There has long been a dispute as to whether the new material is added by "intussusception" throughout the thickness of the wall, or whether it is added to the interior surface, i.e. by "apposition". The bulk of the evidence now supports the second view, at least for many types of cell, but the possibility that there is also some intussusception cannot be excluded. Before we can consider the problem of wall growth further, however, it is necessary to consider the structure of the wall.

Electron microscopic studies have shown that the main structural element of the wall in higher plants consists of a framework of cellulose *microfibrils* (Fig. 1.4), which are somewhat flattened in cross-section, having a width of 10–30 nm,† a thickness of 5–10 nm and a length of at least 60 nm. The cellulose of the microfibrils is mainly present in a crystalline state, i.e. the molecules are regularly arranged in a lattice, while the remainder is semi- or para-crystalline. The microfibrils are embedded in a continuous matrix, consisting mainly of the so called “*hemicelluloses*” (non-cellulosic polysaccharides, composed mainly of residues of the pentoses, arabinose and xylose, and the hexoses, glucose, galactose and mannose) and “*pectins*”, which contain a high proportion of galacturonic acid residues. The matrix also contains low amounts of proteins and lipids. (Further details of the composition of cell walls are given in Chapter 4, pp. 84–85.)

Growth of the wall involves the yielding of the wall to the stress generated by turgor pressure. During the extension of the walls the microfibrils become reoriented. In a typical parenchymatous cell undergoing elongation, the microfibrils are at first oriented in a transverse direction (i.e. at right angles to the long axis of the cell), but as the wall becomes stretched they may be arranged predominantly along the longitudinal axis. During growth, however, new transverse microfibrils are added to the inside of the wall, so that in a cross-section of the wall we find a gradual transition from transversely to longitudinally oriented microfibrils, in passing from the inside to the outside (Fig. 1.5).

The increased plasticity of the cell wall during vacuolation, referred to earlier, must indicate that the various types of chemical bond which link the different wall components must be broken during wall growth, possibly as the result of the activities of hydrolytic enzymes (p. 87).

In many types of cells, growth occurs fairly uniformly over the whole wall, giving the so-called “multi-net” pattern of growth. In other cases, as in root hairs and pollen tubes, the cell may extend by “tip growth”; in such cases it is found that in the growing tip region of a cell the microfibrils are oriented in a random fashion (Fig. 1.4), but during the process of wall stretching they become predominantly oriented in the direction of the cell axis. (Figure 1.4 relates to the primary wall of the alga, *Valonia*, but a similar structure is found in the tip region of cells which show tip growth.)

It is not known what determines the initial transverse arrangement of the microfibrils, but it is found that they usually lie parallel to certain *microtubules* which, as the name suggests, are elongated cylindrical structures of diameter 23–27 nm, found in the boundary layers of the cytoplasm (Fig. 1.6). Moreover, treatment with colchicine, which disrupts the microtubules, also disorganizes the arrangement of the microfibrils, but does not prevent their deposition. Thus, the orientation of the microtubules may, in some way not yet understood, determine that of the microfibrils of the wall.

The Golgi bodies also appear to play a role in cell wall synthesis, since they are conspicuous in regions of active wall synthesis, especially during the development of the cell plate following division (see below). Moreover, vesicles formed by the Golgi bodies have been

† Nanometers = $1 \text{ m} \times 10^{-9}$.

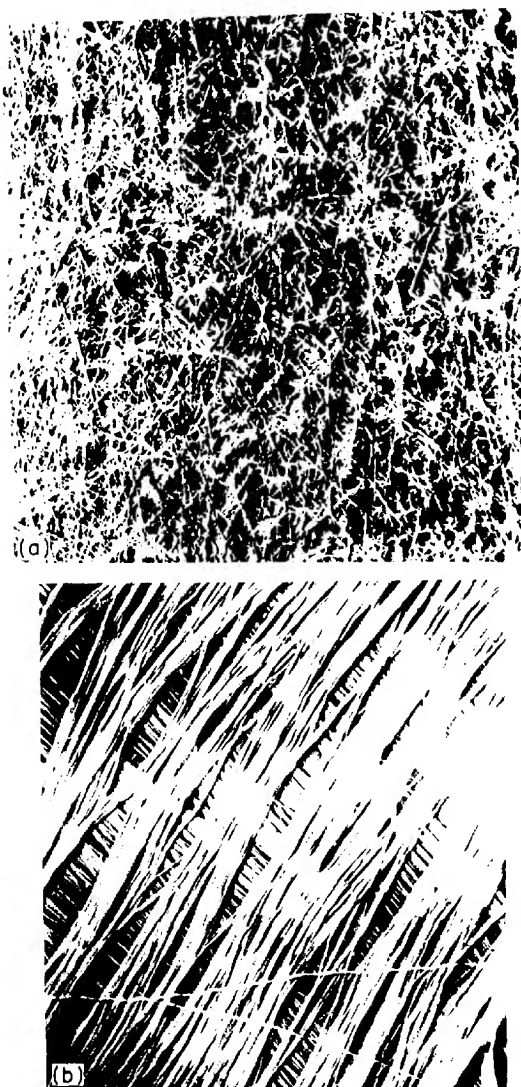


FIG. 1.4. (a) Electron micrograph showing structure of the primary wall of *Valonia* $\times 8000$. (b) As above, but of the secondary wall $\times 7000$. (From F. C. Steward and K. Mühlethaler,

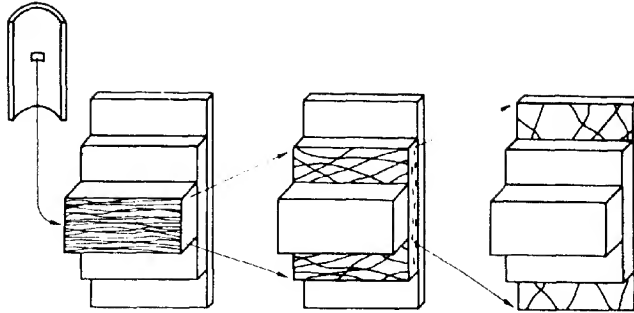


FIG. 1.5. The multinet concept of cell wall growth showing (left to right) re-orientation of microfibrils as successive stages of wall extension. (From P. A. Roelofs, *Adv. Bot. Res.* **2**, 69-150.)

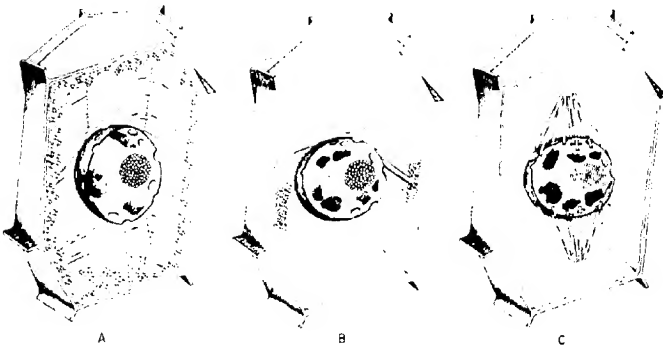


FIG. 1.6. Diagram of changes in microtubules at cell division. (Prints supplied by Dr. Myron Ledbetter, reproduced from *Symposium Int. Soc. Cell. Biol.* **6**, 1967.)

shown to contain polysaccharide material. It is possible that the Golgi bodies are responsible for the deposition of the hemicelluloses and pectins of the matrix of the cell wall.

THE FORMATION OF NEW CELL WALLS

Cell division involves the formation of a new cell wall between the two daughter cells. The process commences with the appearance of large numbers of vesicles in the plane of the equator of the spindle. These vesicles are apparently formed by Golgi bodies, and may contain polysaccharides from which the first stage of the new wall, known as cell plate, is

formed by coalescence of the vesicles. The cell plate forms first in the centre of the cell and its edge extends outwards, apparently by the addition of vesicles from Golgi bodies which lie on the periphery of the plate, until it joins up with the lateral walls (Fig. 1.3.)

Since the new wall is formed at the equator of the spindle, the plane of the wall is determined by the orientation of the spindle. Recent electron microscopic studies on dividing cells in the roots and coleoptiles of wheat suggest that the orientation of the future spindle is determined by certain changes occurring in the cytoplasm before the nuclei enter prophase. In resting cells the microtubules lie in the outer cytoplasm, just inside the plasmalemma. In cells which are about to undergo division, but in which the nucleus has not yet entered prophase, the "wall microtubules" just described disappear and a band consisting of a large number of microtubules appears in the outer cytoplasm near the longitudinal walls, and at right angles to the axis of the cell. The band appears to run right round the wall surface, in the mid-region of the cell (Fig. 1.6).

In cells which divide equally, the plane and position of the preprophase band of microtubules coincide with those of the future new wall, but certain of the epidermal cells of the roots of *Phleum pratense* divide unequally by the asymmetric positioning of the new cross wall (p. 317), and in such cells the band of microtubules still appears in the mid-region of the cell. It would appear, therefore, that the band is not primarily concerned with the location of the cell plate, but with the orientation of the spindle, though how this is achieved is not known.

MEASUREMENT OF GROWTH

So far we have been concerned largely with qualitative and descriptive aspects of plant growth. It is also important to study growth quantitatively, however, and for this purpose we need methods for measuring growth.

As we have already seen, in the final analysis, we might say that growth involves an increase in the amount of living material. However, it is not always easy to measure this increase in living material without destroying the organism in the process. Moreover, if we simply include the protoplast material (cytoplasm and nucleus), we shall leave out increase in such materials as cell walls which form an integral part of the plant body.

It is possible to adopt a different approach to the problem. Since growth essentially involves increase in cell number, we may use this criterion as a measure of growth. Thus, we can measure the growth of a colony of unicellular organisms by counting the increase in the number of individual cells.

In multicellular organisms, such as higher plants, growth still involves large increases in cell number, but it clearly is inconvenient, if not impossible, to measure such increases. However, this increase in cell number, which is accompanied by cell growth, leads to an increase in size and in the case of a root or an unbranched shoot it may be convenient to measure simply the increase in length or height over a given interval of time. This method

is not usually appropriate for a complex root or shoot system, however. Since, if we are studying the growth of a whole plant, we are concerned with the increase in total new tissue formed, it is frequently most appropriate to study changes in the *dry weight* of the plant, which will reflect the actual amount of new organic material synthesized by the plant. However, even a change in dry weight is not always a satisfactory measure of growth, since plant tissues may increase in dry weight due to the accumulation of reserve materials, such as starch and fat, although they may not be growing. Conversely, a germinating seed may show an overall loss in dry weight, due to the utilization of reserves in respiration, although there is no doubt that it is growing.

COLONY GROWTH IN MICRO-ORGANISMS

Before considering the growth curves of higher plants, it is useful to study the growth of a colony of unicellular organisms, such as bacteria or yeast, which multiply by division or "budding", or of a multicellular organism, such as duckweed (*Lemma*), which similarly multiplies by a form of budding.

Consider the growth of a colony of bacteria maintained under constant nutritional and environmental conditions, so that there is a constant rate of cell division. Assume also that the cells divide synchronously, i.e. all cells in the colony divide simultaneously. (Synchronous division can be achieved with cultures of certain organisms.) If the initial number of cells in the colony is n_0 , and n the number of cells after a given number of divisions, then,

at the end of the 1st generation $n = n_0 \times 2$,

at the end of the 2nd generation $n = n_0 \times 2 \times 2$,

at the end of the x th generation $n = n_0 \times 2^x$.

This latter relation indicates that the number of cells in the colony is increasing by geometric progression or "exponentially", i.e. at an ever-increasing rate, and if we plot n against the number of generations, we obtain a curve of the form seen in Fig. 1.7A.

We can rewrite the equation $n = n_0 \times 2^x$ as:

$$\log n = \log n_0 + x \log 2. \quad (1)$$

It will be seen that we have an equation expressing the relation between the number of cells in the colony n , and x , the number of generations which have occurred, but normally we require the relation between n and t , the time. Now, if t is the time taken for x generations, and the time of one generation (i.e. time between two successive divisions) is g , then $x = t/g$.

Substituting in equation (1), we get

$$\log n = \log n_0 + t/g \log 2.$$

Now,

$$\log 2/g \text{ is a constant } (k).$$

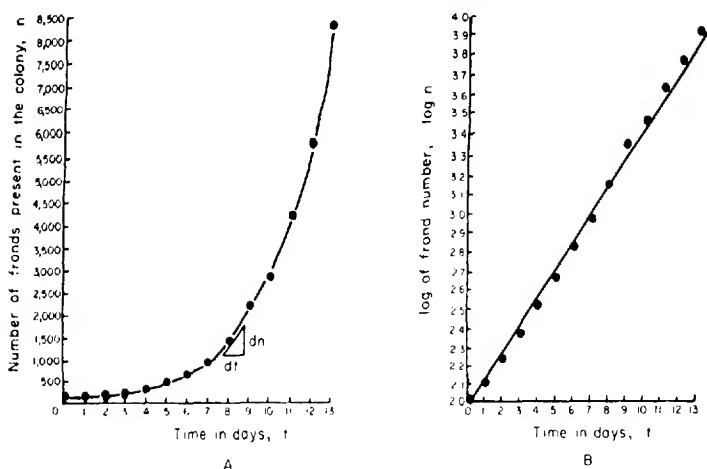


FIG. 1.7. A. Growth curve for a colony of duckweed (*Lemna*) budding synchronously at a constant rate. The initial number of fronds is assumed to be 10. B. Growth of a colony of *Lemna* in culture. Linear relationship between logarithm of frond number ($\log n$) and time. (Data from E. Ashby and T. A. Oxley, *Ann. Bot.* **49**, 309, 1935.)

Therefore,

$$\log n = \log n_0 + kt. \quad (2)$$

Now equation (2) is a linear equation of the form $y = a + bx$, where $\log n_0$ corresponds to a , and k corresponds to b . Hence if we plot the \log of the number of cells present in the colony after different times, against t , we should obtain a straight line. This relation is, in fact, found to hold in practice for various organisms growing under constant conditions, whether they multiply by fission, as in bacteria, or by budding, as in yeast and duckweed (*Lemna*) (Fig. 1.7B). A colony growing in this manner is said to be increasing "logarithmically" or "exponentially".

If we consider the type of curve shown in Fig. 1.7A, giving the increase in the number of *Lemna* "fronds" in the colony, then the *growth rate* of the colony at any time is given by the increase (dn) in the number of cells over a short interval of time, dt , or we can say, growth rate $\propto dn/dt$.

The value dn/dt represents the *slope* of the curve at any given time, t , and it will be seen from the graph that the value of the slope increases progressively with time. If all cells are dividing at the same rate, r , clearly at any time, t , the rate of growth of the colony is proportional to the number of cells present, i.e. $dn/dt \propto n$. Thus, although the rate of cell division (r) remains constant, the *absolute growth-rate* of the colony as a whole does not, since as time goes on the number of cells present in the colony increases. The value of r is

clearly given by dn/dt (the increase in cell number over a given short interval of time), divided by the number of cells so dividing, i.e.

$$r = \frac{dn}{dt} \cdot \frac{1}{n}$$

and this value is known as the *relative growth rate* of the colony. Thus, for a colony showing this type of growth, the absolute growth rate increases with time, but the relative growth rate remains constant.

It has been shown above (equation (2)) that

$$\log n = \log n_0 + kt.$$

This equation can also be written in the form

$$n = n_0 e^{kt} \quad (3)$$

where $e =$ the base of natural logarithms (2.7182).

In practice, unrestricted growth of a colony can never proceed indefinitely and some limiting factor, such as deficiency of nutrients, must always lead to a decline in growth rate sooner or later. Under cultural conditions in a flask or tube, for example, the food supply will ultimately be exhausted and growth will finally cease. Instead of the typical "exponential" growth curve for cell number, we obtain a "sigmoid" type of curve (Fig. 1.8 *left*), in which the growth rate increases up to the point of inflection and then declines gradually to zero. When $\log n$ is plotted against time, growth follows a straight line initially, but later declines (Fig. 1.8 *right*). In addition to the exhaustion of some food factor, growth in colonies may also be limited by some toxic substance which is formed during growth. The production of such "staling factors" often occurs in cultures of bacteria, fungi, *Chlorella*, etc.

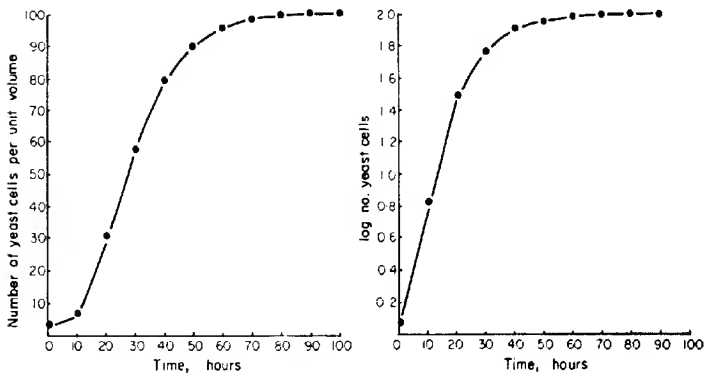


FIG. 1.8. Growth of a colony of yeast growing in a constant volume of culture solution. *Left*: Sigmoid growth curve obtained when number of cells (n) is plotted against time. *Right*: Plot of logarithm of cells ($\log n$) against time. (Data from O. W. Richards, *Ann. Bot.* **42**, 271, 1928.)

GROWTH OF MULTICELLULAR ORGANISMS

1. The Exponential Phase

The sigmoid type of growth curve observed for colonies of unicellular organisms is characteristic also of the growth of individual multicellular plants. This is true not only for the whole plant (Fig. 1.10), but also for individual organs, such as leaves (Fig. 1.9) or internodes. Initially the organism is increasing in size (or weight) by geometrical progression or exponentially. V. H. Blackman (1919) showed that during this initial phase the growth of seedlings follows a "Compound Interest Law" fairly closely and is given by the equation

$$W = W_0 e^{rt} \quad (4)$$

where W = weight of plant after time t ,

W_0 = initial weight of plant,

r = percentage (or proportional) rate of increase,†

e = exponential coefficient (=2.7182).

This equation is clearly exactly comparable with equation (3) above for colony growth and may be derived in a precisely similar manner. From equation (4) we may write

$$\log W = \log W_0 + rt \log e.$$

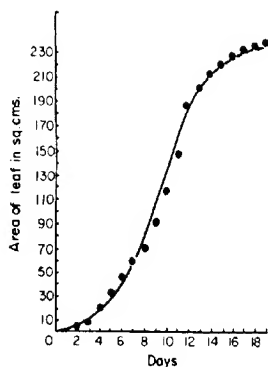


FIG. 1.9. Sigmoid growth curve of leaf of cucumber (*Cucumis sativa*). (From F. G. Gregory, *Ann. Bot.* 35, 93, 1921.)

† The same symbol, r , has been used here as for the rate of cell division in a colony of bacteria, described above, since in both cases r represents the relative growth rate, whether measured by rate of cell division or increase in dry weight.

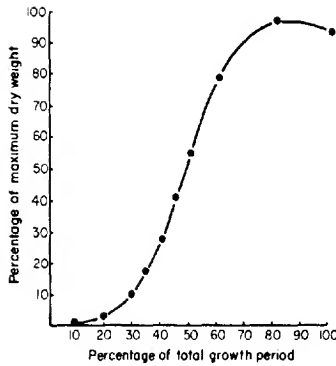


FIG. 1.10. Increase in dry weight of barley plants during growing season. The points on the curves were derived from smoothed curves drawn from the transformed original data. (From F. G. Gregory, *Ann. Bot.* 40, 1, 1926.)

This equation is again of the form $y = a + bx$. This means that we should obtain a straight line when we plot log weight against t (at least for the initial phase of growth, which we are now considering) and this has, in fact, been demonstrated in a number of cases.

From equation (4), it is clear that the final weight attained will depend upon (1) the initial weight, (2) the rate of "interest" (r), (3) the time. The rate of "interest" represents the efficiency of the plant as a producer of new material and was called by Blackman the *efficiency index* of dry weight production. A small difference in the efficiency index between two plants will soon make a marked difference in the total yield, and the difference will increase with the lengthening of the period of growth.

It should be noted that the efficiency index is merely a different method of expressing the *relative growth rate* ($dW/W \cdot dt$) as described for colony growth. Whereas the efficiency index (or relative-growth rate) remains constant through the exponential growth phase, the *absolute* increments per unit time increase progressively. The absolute growth increment over a time interval dt is clearly

$$\left(W \times \frac{r}{100} \right) \cdot dt.$$

Thus, the absolute growth rate at any given time is proportional to the size of the plant at that time.

The physiological basis of this latter conclusion is easily understood, for when photosynthesis has become active in a young seedling, the power of the plant to synthesize new material (and hence increase in dry weight) is clearly dependent upon its leaf area. Therefore, as the plant grows and increases its leaf-area, the rate at which new material is assimilated will increase proportionately.

2. Later Phases of Growth

Just as the growth rate (dw/dt) of a bacterial colony ultimately falls off with time due to the exhaustion of nutrients or the accumulation of toxic products, so the growth rate of a multicellular organism decreases gradually, resulting in a sigmoid growth curve. The absolute growth rate (dW/dt) is clearly given by the slope of the growth curve at any time t and if we plot the changes in this growth rate with time we get a curve of the type shown in Fig. 1.11A. We see that the growth rate attains a maximum (corresponding to the point of inflection of the S-shaped growth curve) and then falls away to zero. If, on the other hand, we plot the relative growth rate ($dW/W \cdot dt$) against time, we frequently get a curve of the type in Fig. 1.11B. It is seen that the relative growth rate (RGR) remains nearly constant at first but later begins to decline. The changes in RGR with time vary a great deal from one species to another and with the conditions under which the plants are growing. Sometimes it is found that the RGR declines steadily from the commencement of growth, so that a true exponential phase does not occur.

The reason for the fall in the RGR is not fully understood and various hypotheses have been suggested. The deficiency of some nutritive factor is clearly not the cause, as it is in colonies of unicellular organisms under artificial conditions. It has been suggested, however, that the reason for the departure from "exponential" growth in a normal plant is that part of the plant material formed during growth gives rise to mechanical, vascular and other tissues which do not directly contribute to further synthesis of new material. The leaves which are the organs most directly concerned in the synthesis of new material thus constitute a diminishing fraction of the total plant weight, i.e. the ratio

$$\frac{\text{Total leaf dry weight } (L)}{\text{Total plant dry weight } (W)},$$

known as the *leaf-weight ratio*, gradually falls.

Now, the relative growth rate

$$(R) = \frac{dw}{dt} \cdot \frac{1}{W}.$$

Multiplying numerator and denominator by L we get

$$R = \frac{dw}{L} \cdot \frac{L}{W} \cdot \frac{1}{dt}.$$

Now $dw/L \cdot dt$ is the rate of increase in dry weight per unit dry weight of leaf, known as the *net assimilation rate* (E_w) and is a measure of the photosynthetic activity, minus losses due to respiration; that is, E_w is a measure of the net efficiency of the plant in the production of dry matter.

It will be seen that we have the following simple relationship between the three parameters:

$$R = E_w \times \frac{L}{W}. \quad (5)$$

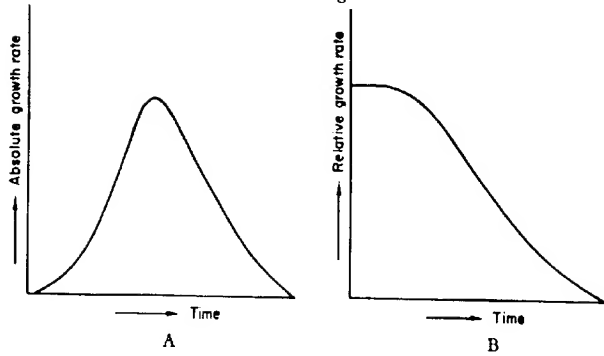


FIG. 1.11. Changes in (A) absolute growth rate dW/dt and (B) relative growth rate $(dW/W \cdot dt)$, where the dry weight changes follow the type of curve shown in Fig. 1.10.

If we assume that E_w does not change appreciably with the age of the plant, then the fall in R with age must primarily be due to the fall in L/W , for the reasons already indicated. However, both E_w and L/W are frequently found to decline during the growth period and thus to contribute to the decline in R .

In the above discussion, we expressed the net assimilation rate (E_w) in terms of the rate of increase in total plant dry weight per unit leaf dry weight, but it is more usual to express photosynthetic efficiency as the rate per unit *area* of leaf. However, the net assimilation rate can easily be expressed on a leaf area basis, by making use of the following relationship:

$$E_w = E_A \times \frac{L_A}{L_w}$$

where E_A is the rate of increase in dry weight per unit leaf area and L_A/L_w is the ratio of leaf area to dry weight, known as the *specific leaf area*.

Thus, equation (5) above can be rewritten as

$$R = E_A \times \frac{L_A}{L_w} \times \frac{L_w}{W}$$

The foregoing simple mathematical relationships have been used to analyse the growth of crops. Thus, the determination of relative growth rates for different crop plants gives us a useful basis for comparing their growth rates. Similarly by determining net assimilation rates and leaf-weight ratios we can obtain some indication of how differences in R arise.

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CHAPTER 2

Patterns of Growth and Differentiation

LEVELS OF DIFFERENTIATION

So far, we have been concerned primarily with growth, rather than differentiation. In the present chapter we shall describe the development of the main organs of the plant and the way in which differentiation arises within the plant.

Now, when we consider the manifold forms of differentiation in the plant it is evident that it occurs at various levels. At the highest level, there is differentiation in the plant body as a whole, as seen in the division into root and shoot. Within the shoot we can observe the differentiation into various organs, such as stems, leaves, buds and flowers, and within each of these organs there is differentiation at the cellular and tissue level. These three levels of differentiation also constitute a series of successive stages in *time*—there is first differentiation into root and shoot in the embryo, and this is followed by the formation of organ primordia, as a result of the activities of the apical meristems. These organ primordia do not at first show differentiation at the cell and tissue level, which occurs during the later stages of their development. We may illustrate the progressive steps of differentiation of the plant body in a diagram (Fig. 2.1), in which it is seen that at various stages there are divergent alternative pathways, leading to successively more specific pathways of differentiation.

The chain of developmental events usually takes place in a very orderly manner, one stage following another in a proper sequence. This orderly sequence of changes suggests that the successive steps are not controlled independently, but that the attainment of one stage exercises some control over the alternative pathways which will be entered at the next step. This idea is far from new, since it was clearly enunciated by Pfeffer in 1903.

Once a particular pathway of development has been entered, the process is usually irreversible; for example, once a flower primordium has been formed, it is not easily converted back to a leafy shoot. Similarly, cortical cells do not normally change directly into vascular elements nor vice versa. Thus, at certain stages organs and tissues become *determined* in their pattern of development. Nevertheless, some tissues do retain the capacity for “de-differentiation”, so that roots may be initiated in stem tissue, and so on. The successive entry into

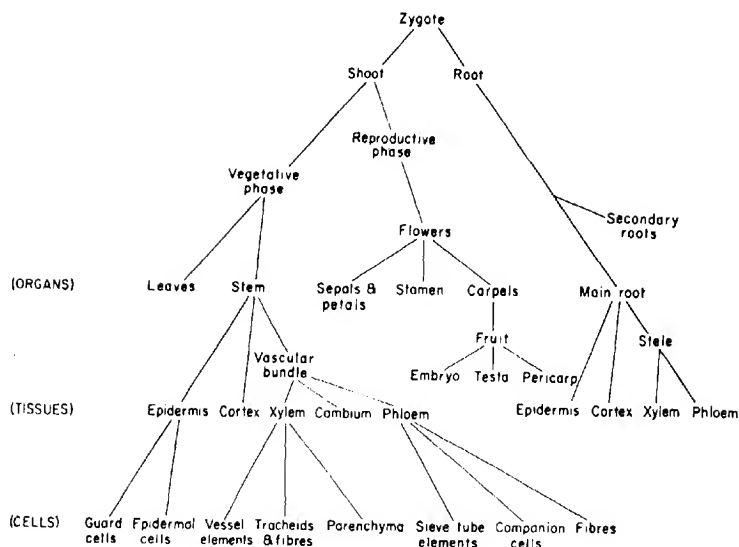


FIG. 2.1. Levels of differentiation during the development of a higher plant.

specific developmental pathways which, once entered, cannot easily be departed from, has been referred to by Waddington as the "canalization of development".

In addition to the first major step in differentiation (viz. the formation of root and shoot), certain other changes occur during the life cycle of seed plants, which must be regarded as aspects of differentiation, of which the most important is the transition to the reproductive phase, which involves a profound change in the structure of the shoot apex (p. 44). We shall see later that in many species the onset of flowering is controlled by environmental factors, but in many other species it appears to be determined more by progressive changes occurring during the development of the plant itself than by environmental factors. Often these progressive physiological changes are reflected in morphological characters, such as leaf shape, in which a gradient up the stem may frequently be seen. These aspects of development will be dealt with later, but it is important to recognize that they represent an aspect of differentiation within the shoot as a whole.

TISSUE AND CELL DIFFERENTIATION

When the development of the main organs has been initiated, with the determination of stem, leaf, root and flower, each of these organs follows a largely independent course of

further development, as shown by the fact that a leaf primordium will undergo full differentiation if isolated and maintained in a septic culture (p. 36). However, in the intact plant development is a continuous process and it is somewhat artificial to draw too sharp a distinction between organ and tissue differentiation. In particular, the differentiation of vascular tissue is continuous throughout the plant and this continuity is no doubt an important factor in achieving co-ordination of development and function in the plant as an integrated whole.

At the cellular level, the term differentiation is sometimes used in two different senses: viz. (1) it may be applied to the development of different specialized types of mature cell within an organ or tissue; or (2) it may be used to refer to the changes which occur during the development of a meristematic cell into a mature cell, usually involving vacuolation and enlargement. In this book we shall use the term in the first sense and we shall use the term *maturation* for the processes which lead to the formation of a mature cell from a meristematic one.

Usually maturation involves cell vacuolation and enlargement, and some aspects of this process have already been described (p. 7). Cells undergoing maturation may show relatively little other change in structure, as in the formation of parenchymatous tissue, or there may be great changes, as in the formation of xylem and phloem tissue. It is the diverging pathways followed by different cells during maturation which result in differentiation.

In addition to the visible changes involved in differentiation, there are also biochemical differences, some of which will be described later (p. 312).

Between the biochemical differences occurring at the molecular level and the structural changes observable with the optical microscope, one might expect to find changes at the ultrastructural level, to be seen under the electron microscope. Studies on various types of tissue have shown that, in general, living differentiated higher plant cells have the normal cell organelles, including mitochondria, Golgi bodies (dictyosomes), plastids and endoplasmic reticulum, but there are certain exceptions, e.g. sieve tubes, in which most of the organelles disintegrate during differentiation. The organelle which shows the most marked differences in various types of tissue is the plastid, the structure of which varies enormously according to whether it occurs in leaf tissue, storage tissue, fruits (as in tomato), or flower parts, such as petals. Mitochondria also vary quite markedly in number and structure in different types of cell and the Golgi bodies show active and quiescent states related to the state of wall growth, secretion and so on. The endoplasmic reticulum varies in abundance and localization in several types of specialized cell, particularly those concerned with secretion.

On the other hand, many of the differences to be observed between various types of tissue relate to the cell wall. Not only do we find very characteristic differentiation of the cell wall in, for example, the various types of xylem cell, but also in living differentiated cells, such as collenchyma, endodermis and stomatal guard cells. These readily observable differences reflect differences detectable at the ultrastructural level. However, an account of the ultrastructural changes in the plastids and cell wall occurring during differentiation is beyond the scope of this book.

Most cells which have attained the mature condition do not normally undergo subsequent cell division. However, some types of mature cell still retain the capacity to divide, as seen in the origin of vascular cambium, cork cambium and adventitious root initials in various types of parenchymatous tissue. Moreover, many types of cell are able to resume cell division in response to wounding (p. 158).

DIFFERENTIATION INTO ROOT AND SHOOT—EMBRYO DEVELOPMENT

The first major step in differentiation occurs at a very early stage in the development of the embryo, with the establishment of a shoot end and a root end.

As an example of embryo development in angiosperms we may consider that of *Capsella bursa-pastoris* (Fig. 2.2). The fertilized zygote is a somewhat elongated cell, which divides transversely to give a smaller *terminal* cell, and a larger *basal* cell. The terminal cell gives rise to most of the future embryo, whereas the basal cell gives rise mainly to the *suspensor*. The terminal cell divides by two successive longitudinal divisions, with the plane of the

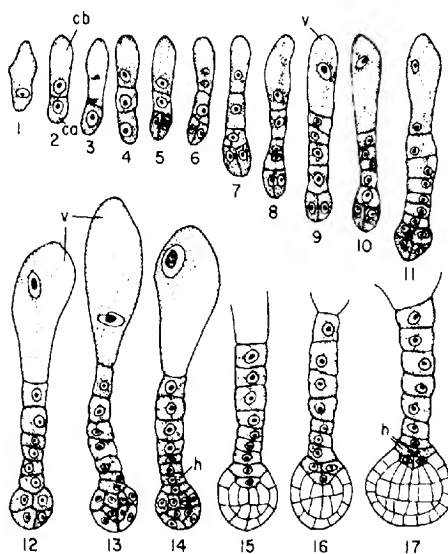


FIG. 2.2. Early stages in the development of the embryo of *Capsella bursa-pastoris*. Note the initial unequal division into a larger basal cell (cb) and smaller terminal cell (ca). The basal cell gives rise to the suspensor. (From A. Fahn, *Plant Anatomy*, Pergamon Press, Oxford, 1967, adapted from Souèges, 1914.)

second division at right angles to that of the first, to give four cells; these cells then each divide transversely to form eight cells, which constitute the octant. Each octant cell divides to form an outer protodermal cell, which gives rise to the future epidermis, and an inner cell. The inner cells continue to divide to form the cotyledons and the hypocotyl.

By several successive transverse divisions the basal cell gives rise to a row of cells which forms the suspensor, the end cell of which enlarges and becomes sac-like. The suspensor cell nearest the embryo undergoes several divisions, to give a group of cells, of which the outer ones form the future root cap and root epidermis, while the inner ones form the remainder of the radicle. The fully developed embryo is formed by further repeated divisions of these various regions.

There is considerable variation from the pattern of development described for *Capsella* among the various groups of angiosperms, but the details do not concern us here. However, whatever the variation in further development, the initial stages have certain features in common, namely that the first division of the zygote gives rise to two unequal cells, and of these, the basal cell is normally the one nearer the micropyle of the ovule, and gives rise to the root end of the embryo, whereas the terminal cell gives rise to the shoot end. Thus, even the very young embryo shows *polarity*, in that it has a shoot end and a root end. Indeed, the egg itself shows differences in the density of cytoplasm between its two ends, suggesting that the first unequal division of the zygote is already predetermined by polarization in the unfertilized egg. We shall discuss the basis of this polarity in more detail later.

After the embryo has become differentiated into root and shoot regions, apical meristems are established and some organs become differentiated, often while the seed is still developing on the parent plant, so that not only cotyledons but also a rudimentary epicotyl and, in grasses, even several leaf primordia, may be present.

SHOOT APICAL MERISTEMS

Although this book is primarily concerned with flowering plants, it is useful to consider, briefly, the patterns of growth in the lower plants. In simple filamentous algae, such as *Spirogyra*, every cell appears to be potentially capable of division and growth and is not localized to particular regions. However, in many algae there is marked localization of growth. Thus, the alga, *Chara*, has a single prominent apical cell, which divides repeatedly, giving a larger outer (distal) cell, which continues to function as the apical cell, and a smaller proximal daughter cell, which proceeds to undergo further division, the resulting cells forming the mature tissue of the thallus.

In the bryophytes and many pteridophytes, also, the shoot grows by a single well-marked apical cell, which normally has the form of an inverted tetrahedron, and it divides so that the three "inner" faces cut off daughter cells in succession, and these latter cells undergo further division to form the tissues of the shoot (p. 320).

The early plant anatomists of the nineteenth century were so impressed with the essential unity of structure in vascular plants that they expected to find single apical cells also in

gymnosperms and angiosperms and indeed described such cells. Later, however, it became apparent that there is no clearly recognizable *single* apical cell in the shoots of higher plants, but two zones may be distinguished in the shoot apical region of flowering plants: (1) the outer *tunica* or mantle, which surrounds and envelops, (2) the inner *corpus* (Fig. 2.3). These zones can be distinguished fairly sharply by the predominant planes of cell division. In the tunica the divisions are predominantly *anticlinal*, i.e. with the axis of the mitotic spindle parallel to the surface, so that the resulting cross-wall separating the two daughter cells is perpendicular to the surface. The corpus, on the other hand, is characterized by the fact that divisions occur in all planes, viz. both *anticlinal* and *periclinal* (i.e. the spindle is perpendicular and the new wall parallel, to the surface). The thickness of the tunica is somewhat variable and it may consist of one, two or more layers of cells, according to the species. Moreover, even within a single species the number of tunica layers may vary according to the age of the plant, the nutrient status and various other conditions.

It should be noted that the tunica-corpus theory is largely descriptive and simply based on what can be observed—it makes no predictions regarding the future destiny of the tunica and corpus cells. The epidermis does, of course, arise from the outer layer of the tunica, but there is considerable variation between different species in the extent to which



FIG. 2.3. Median section through the shoot apical meristem of *Alternanthera philoxeroides*, showing the two-layered tunica overlying the central corpus. (From E. G. Cutter, *Phytomorph.* **17**, 437, 1967.)

the two layers may contribute to the origin of leaves and buds (see p. 29). However, although we should not regard the tunica and corpus as rigidly and permanently demarcated, it is possible to show that in some species the outer tunica layers remain remarkably distinct from the deeper tissue for long periods. One method by which this has been demonstrated is to induce polyploidy in the cells of the shoot apices of *Datura* and of maize by treatment with colchicine. The tetraploid cells so formed can be recognized by their larger nuclei. If tetraploid cells are formed in the outer layer of the tunica, for example, all the resulting daughter cells will show large nuclei and it can be seen that such cells are strictly limited to a single layer of the tunica (Fig. 2.4), indicating that periclinal divisions are very rare and that each layer of the tunica retains its identity remarkably constantly. Each of the layers of the tunica arises from a set of initial cells at the shoot apex, and the corpus apparently arises from its own set of initials, although there is some difference of interpretation on this point.

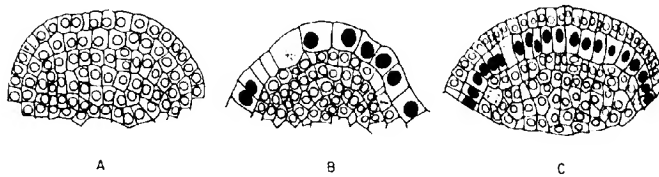


FIG. 2.4. Longitudinal sections of the shoot apex of *Datura*. A. Normal apex, with all cells diploid. B. Apex after treatment with colchicine, showing polyploid nuclei in outer tunica layer. C. After treatment with colchicine, with polyploid nuclei in inner tunica layer. (Adapted from S. Satina, A. F. Blakeslee and A. G. Avery, *Amer. J. Bot.* **27**, 895, 1940.)

The distinction between tunica and corpus is even less definite in gymnosperm shoot apices, and although the divisions in the outer layers of *Pinus*, for example, are predominantly anticlinal, there are also quite frequent periclinal divisions as well.

In addition to the tunica and corpus and their initials, it is possible to recognize a number of other zones in the apices of some species. One simple form of zonation is shown in Figs. 2.4 and 2.5. At the summit of the apical dome there is a group of initial cells which divide mainly anticlinally, and so give rise to the tunica, but the lower layers may also divide periclinally. Below these initial cells there is a group of larger cells, known as the *central mother cells*, which appear to have a low rate of division. Divisions at the boundary of the central mother zone give rise to (1) a zone of actively dividing cells which form the flanks of the apex and is sometimes called the *flank* or *peripheral meristem*; and (2) a zone of cells which divide mainly along the shoot axis and give rise to the longitudinal rows of cells of the cortex and pith and hence is sometimes called the *rib meristem*. The distinction between these zones, which have been given various names by different workers, is somewhat arbitrary, since their boundaries are ill defined and grade into each other. Moreover, there is considerable variation between species, both with respect to the general shape of the apical

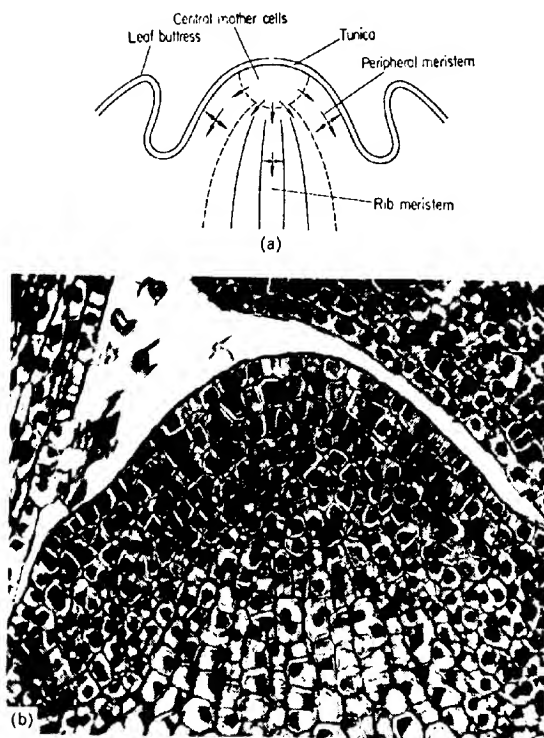


FIG. 2.5. A. Generalized diagram to illustrate zonation in the shoot apical region of a flowering plant. B. Median section through the vegetative apex of *Chrysanthemum morifolium*, in which the various zones shown in A may be distinguished. (From R. A. Popham, *Amer. J. Bot.* 37, 476, 1950.)

region and to the number of zones which can be recognized within it, and yet the apices of all these species produce similar end products, viz. stem, leaves and buds. It would seem, therefore, that not much morphogenetic significance can be attached to the various zones which can be recognized in the shoot apex.

THE INITIATION OF LEAVES AND BUDS

A leaf originates from the periclinal divisions of a group of cells on the flank of the apex (Fig. 2.10). As a result of these divisions in a localized area, a small protuberance (Figs. 2.9

and 2.10) is formed and gives rise to the future leaf primordium. The number of layers involved in these initial divisions varies considerably in different species. In many grasses, the peridental divisions commence in the outermost layer of the tunica, and in the layer below. In other monocotyledons and in dicotyledons, periclinal divisions do not take place in the outermost layer, but in the layers below it. Thus, the extent to which the tunica and corpus are involved in the initiation of the primordium varies greatly. In many species the initial divisions involve both the tunica and the corpus, while in others they may occur in only one or other of these layers. Variations may occur even within a single species.

Lateral buds usually appear somewhat later than leaf primordia, in the sequence of developmental changes seen at the shoot apex. Buds arise in the outer layers of the stem tissues, as a result of cell divisions which may be predominantly anticlinal in the outer layers, or both anticlinal and periclinal in the deeper layers. As with leaves, there is considerable variation between species in the extent to which tunica and corpus are involved in these cell divisions giving rise to lateral buds. As a result of these cell divisions the bud emerges as a protuberance and it soon develops an apical structure similar to that of the main shoot apex for that species.

THE SITING OF LEAF PRIMORDIA

The siting of leaf primordia at the shoot apex is a rather precisely regulated process, although there are considerable differences from one species to another in the pattern of arrangement of leaf primordia. In considering this problem it has to be remembered that the apex is growing continuously and that as it does so the older leaf primordia are left behind on the flanks of the apex and they steadily increase in size as they do so (Fig. 2.6). As the apex grows, new leaf primordia are being continuously initiated above the existing ones.

The siting of leaf primordia at the apex determines the arrangement of leaves on the mature shoot, for which the term *phyllotaxis* is used. The most common type of leaf

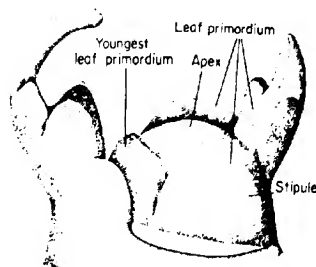


FIG. 2.6. Vegetative shoot apex of *Vitis*. (From A. Fahn, *Plant Anatomy*, Pergamon Press, Oxford, 1967.)

arrangement is *spiral* phyllotaxis, in which it can be seen that a line drawn through successively older leaf primordia at the shoot apex forms a spiral, known as the *genetic* or *developmental spiral* (Fig. 2.7). The mathematical treatment of spiral phyllotaxis has interested botanists for more than a century, but here we shall consider the problem only so far as it relates to the siting of primordia at the shoot apex.

The fern apex provides very convenient material for studying phyllotaxis, since it is relatively flat and the primordia are well spaced out (Fig. 2.8). In considering this problem it is useful to use a system of nomenclature in which the youngest primordium is called P_1 and the successively older primordia, P_2 , P_3 , etc. The next primordium to arise is referred to as I_1 (i.e. Initial₁) and the successively younger primordia as I_2 , I_3 , etc. If we draw radii from the centre of the apex to two successive leaf primordia, it is found that the angle of divergence between the two lines varies from one species to another, but where there are numerous primordia at the apex, it is found to approach a "limiting" value of 137.5° . It is also found that in many species the radial distance of successive primordia increases in geometric progression, indicating that there is a corresponding increase in the rate of expansion of the apex with distance from the centre. Thus, we can reproduce the spiral phyllotaxis seen at a shoot apex by marking points consecutively around a centre at a constant divergence of 137.5° and radially at a distance from the centre which increases in geometric progression. If we then join the successive points, we shall obtain a genetic spiral.

What determines that the next primordium will be formed at a position which will cause an angular divergence from the preceding primordium of approximately 137.5° ? There has been a great deal of controversy on this question, but at present there are two main theories to account for the observed facts, which may be referred to as the "Repulsion Theory" and the "Available Space Theory", respectively. Both theories postulate that the positions in which the leaf primordia arise are determined by the positions of older primordia. According to the Available Space Theory (first put forward by Hofmeister in

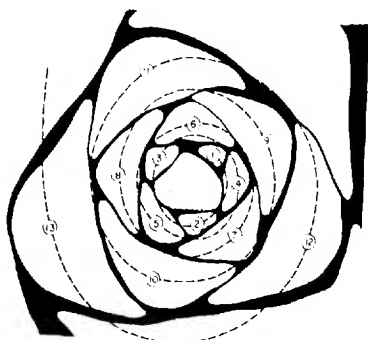


FIG. 2.7. Diagram to illustrate spiral phyllotaxis as seen in cross-section through shoot apical region of *Saxifraga* (genetic spiral shown by broken line). (From F. Clowes, *Apical Meristems*, Blackwell Scientific Publications, Oxford, 1961.)

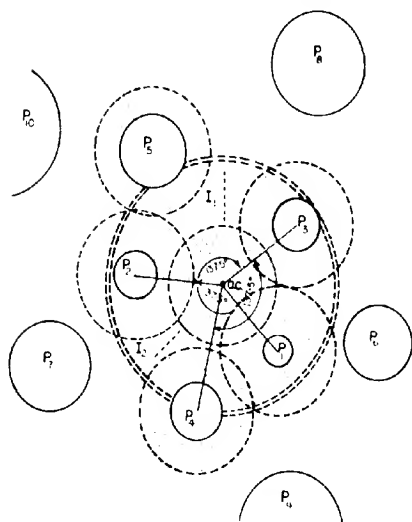


FIG. 2.8. The growth centre and field concept as it may apply to the apex of *Dryopteris*. The apex is seen from above. *ac*, apical cell; P_1, P_2, P_3, P_4 , etc., leaf primordia in order of increasing age; I_1, I_2 , the next primordia (as yet invisible) to be formed in that order. The large double-dashed circular line indicates the approximate limit of the apical cone. The subapical region lies outside the double-dashed circle. The hypothetical inhibitory-fields around the several growth centres are indicated by stippling. (Adapted from C. W. Wardlaw, *Growth*, 13, Suppl. 93, 1949.)

1865) a certain minimum space between existing primordia and the centre of the apex is necessary before a new primordium can arise. As the apex grows, the spaces between existing primordia increase in size and it is postulated that the next primordium (I_1) arises between P_2 and P_3 because this is the first area to reach the necessary minimum size. The next primordium after this (I_2) will arise between P_2 and P_4 , and so on.

The Repulsion Theory was put forward by Schoute (1913) who postulated that the centre of a leaf primordium is determined first, and that a specific substance is produced which inhibits the formation of others in the immediate vicinity, so that new primordia again arise in the gaps between older ones, where they will presumably be outside the inhibitory fields of the neighbouring primordia (Fig. 2.8). Inhibition of new centres by the main apex was also postulated, so that new primordia are prevented from forming within a minimum distance from the summit of the apex. As we shall see later (p. 335), there is, indeed, evidence for the existence of mutual inhibition between growth centres.

These two hypotheses are not necessarily exclusive, since the "available space" may not be determined simply by the superficial area between adjacent primordia, but by freedom from their inhibitory influence. Both theories postulate that phyllotaxis is determined

primarily by the geometry of the shoot apex. It can be shown, from purely geometrical considerations, that I_1 will arise at a divergence of 137.5° from P_1 , if it occurs at a point which divides the angle between P_2 and P_3 in the inverse ratio of their respective ages; that is, the new primordium will not be equidistant from the neighbouring primordia, but will be displaced towards the older (and larger) of these (Fig. 2.8). It is not clear what is the significance of this fact, but it is consistent with the hypothesis that the neighbouring primordia play an important part in determining the position of a new primordium.

Attempts have been made to test the "available space" and "repulsion" theories by surgical experiments on the shoot apex. For example, in experiments with *Lupinus albus*, R. and M. Snow made radial cuts in the area at which I_2 would be expected to arise, thereby reducing the space available, and it was found that no primordium developed in this space, presumably because it was reduced below the minimum area necessary for primordium development. On the other hand, Wardlaw isolated I_1 of fern apices by two radial cuts and found that it then grew *more rapidly* than normally (Fig. 2.11B), suggesting that it had thereby been released from the inhibitory effects of neighbouring primordia. Although these and other surgical experiments have produced interesting results, they have not given decisive evidence in favour of either of the two theories.

DEVELOPMENT OF THE LEAF

The overall development of the leaf may be divided into the following steps: (1) formation of the foliar buttress, (2) formation of the leaf axis, and (3) formation of the lamina. The following account is based upon the development of the tobacco leaf.

As we have seen (p. 29), a small protuberance arises on the surface of the flanks of the meristem (Fig. 2.10) by periclinal divisions in the surface layers. Certain cells towards the centre of this *foliar buttress* now begin to divide actively and a small finger-like protuberance emerges from the buttress (Fig. 2.9). This protuberance proceeds to grow in size by the activities of apical cells until it is about 1 mm long. At this stage the leaf primordium consists of little more than the future axis (midrib and petiole) of the leaf. Soon, however, certain cells on its flanks begin to grow, so that it now acquires a more flattened appearance in cross-section. These dividing cells on the flanks of the mid-rib constitute the *marginal meristems* and they give rise to the future lamina of the leaf. The tip growth of the mid-rib ceases fairly early, when the total leaf length is 2–3 mm and further growth is more generally dispersed.

The marginal meristems consist of superficial *marginal initial* cells, together with the underlying *submarginal initials* (Figs. 2.9, 2.10). In dicotyledons, the marginal initials normally divide only anticlinally, to give rise to the surface layers of the leaf, whereas the submarginal initials form the inner tissues of the future leaf. As a result of these activities of the marginal and submarginal initials a definite number of layers is established in the young lamina and under any given set of environmental conditions this number remains rather constant over most of the leaf, throughout its future development. The relative constancy

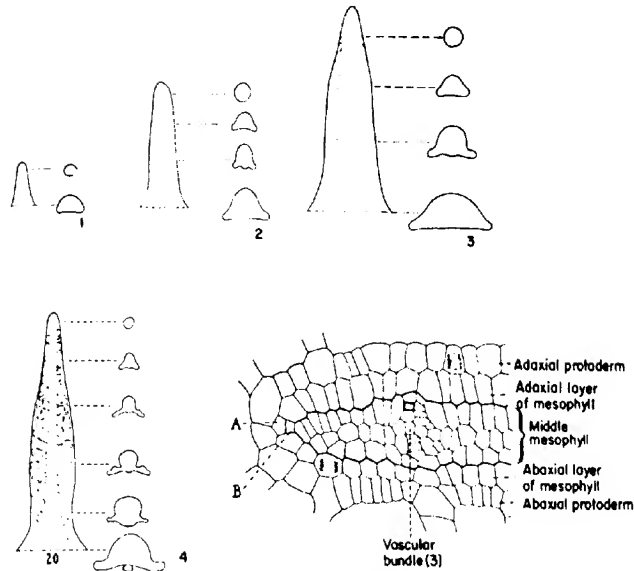


FIG. 2.9. Diagrams of longitudinal and cross-sections of leaf primordia of *Nicotiana tabacum* at different ontogenetic stages. 1. A young, more or less cone-shaped primordium. 2. Primordium in which the narrow margins, from which the lamina will develop, can be seen. 3. Primordium in which the beginning of development of the main lateral veins can be seen. 4. Primordium 5 mm long in which the early development of the provascular system can be seen. 5. Cross-section of the marginal region of tobacco leaf showing submarginal initial and the origin of the mesophyll and a vascular strand. A, B, submarginal initials. (From G. S. Avery, *Amer. J. Bot.* **20**, 565, 1933.)

in the number of layers arises from the fact that the cells continue to divide mainly anticlinally, i.e. at right angles to the surface of the leaf, so that there is a steady increase in area but not in thickness.

The different layers of the leaf are found to cease cell division at different stages. The cells of the upper epidermis usually cease dividing first, when the tobacco leaf is only 6–7 cm long, i.e. when it is only one-fifth to one-sixth of its final size. However, these epidermal cells continue to increase in size until the whole leaf ceases to enlarge. On the other hand, the palisade cells continue to divide and so give rise to a closely packed layer of cells which keeps pace in its growth with the upper epidermal cells. They cease dividing and enlarging shortly before the epidermal cells do so, so that to keep pace with the latter they are pulled apart slightly in the final stages, thus giving rise to the intercellular spaces between the mature palisade cells. The cells of the future spongy mesophyll cease dividing and enlarging

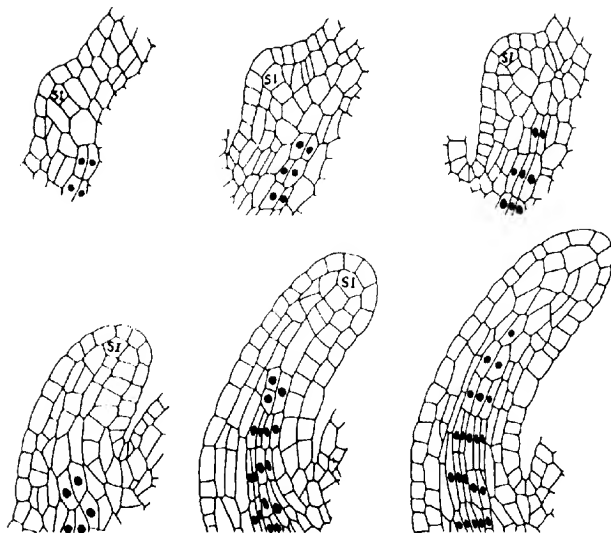


FIG. 2.10. Early stages in development of leaf primordium in flax (*Linum*), showing development of procambial strand (cells with nuclei indicated). S.I., submarginal initial cell. (From G. Girolami, *Amer. J. Bot.* **41**, 264, 1954.)

earlier than the cells of the palisade, so that the cells become pulled apart more in the final stages, giving rise to larger intercellular spaces than in the palisade layers and to a more irregular arrangement of the cells. This study of the origin of the different layers of the mesophyll of the leaf provides a good example of how a study of the morphology of development can give a better understanding of the way in which the mature structure of an organ arises.

The first procambium is formed towards the base of the developing leaf primordium at a very early stage (Fig. 2.10). This first procambium forms the future mid-vein and it develops both outwards towards the tip (acropetally) and downward (basipetally) to link up with the procambium of the stem (Fig. 2.14). The first vascular elements to appear are those of the protophloem, followed later by the protoxylem. Shortly after, the lateral meristems start to form the lamina, and the first signs of the lateral veins appear when the primordium is about 1.55 mm long. The connecting veins soon appear, towards the tip (Fig. 2.9).

Complete normal leaf development depends upon exposure to light and is one of the important aspects of "photomorphogenesis" (p. 188). Hormones, especially auxin and cytokinin, also appear to play an important part in leaf growth (p. 116).

LEAF DETERMINATION

The differentiation of the shoot into leaves, buds and stems commences, as we have seen, with the formation of leaf and bud primordia at the shoot apex.

The early stages of development are very similar in both leaves and buds, but whereas the leaf primordium very early assumes a dorsiventrality (i.e. it becomes flattened, with upper and lower surfaces), the bud remains radially symmetrical. Moreover, whereas the apical meristematic cells of the leaf primordium cease to be active at an early stage and the development of the leaf is determinate, the bud primordium develops a typical shoot apical meristem which shows indeterminate growth.

Although we know from its position and sequence that a given primordium is normally destined to become a bud or a leaf, a very young primordium is not yet irreversibly determined to become one or the other. Indeed, in the early stages of their development, the primordia which, by their position, would normally give rise to leaves, may be converted into buds by certain surgical treatments. These techniques were first developed by R. and M. Snow and have since been used extensively by others, especially by Wardlaw and his associates, using the fern apex. The interconvertibility of leaf and bud primordia has been demonstrated in *Dryopteris dilatata*, by making a deep tangential cut between the shoot apical cell and a very young presumptive leaf primordium, as a result of which it develops into a radially symmetrical bud instead of a dorsiventral leaf (Fig. 2.11A). This conversion of a leaf into a bud can only be effected with very young primordia. When a leaf primordium is a little older it remains as a leaf when such surgical treatment is applied. Thus, the primordia produced on the shoot apex are at first uncommitted, and they only become determined as leaves at a later stage.

A different approach to the problem of leaf determination has been made by Steeves

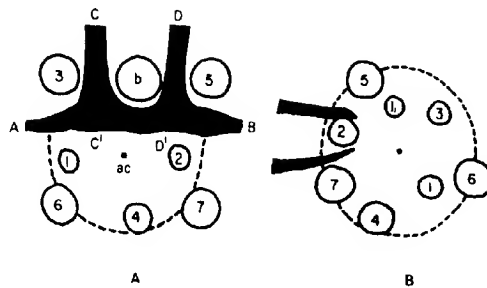


FIG. 2.11. A. Shoot apex of *Dryopteris* in which the 1₁ position was isolated from the apical cells (ac) by a tangential incision (AB) and from leaf primordia 3 and 5 by radial incisions (CC' and DD'). A bud has been formed in what was normally a leaf position. B. Shoot apex in which leaf primordium, P₂, has been isolated from P₅ and P₇ by radial incisions: P₂ grows rapidly and is soon larger than older primordia. (From C. W. Wardlaw, (A) *Proc. Linn. Soc.* **162**, 13, 1950-1; (B) *Growth*, **13**, Suppl'. 93, 1949.)

using sterile culture methods. When primordia are removed from the apex of the fern *Osmunda cinnamomea* and placed on a sterile nutrient medium they are able to undergo further growth and development (Fig. 2.12). When the youngest primordia, P_1 - P_5 , were so tested they developed not as leaves but as shoots which eventually became rooted plants. Progressively older primordia showed an increasing tendency to develop as leaves, and P_{10} always developed as a dorsiventral leaf. From these experiments, Steeves concluded that leaf primordia of *Osmunda* are not irreversibly determined from their inception, but undergo a relatively long period of development during which they remain undetermined. Determination is gradually imposed on the primordium.

Once determination has occurred, the future development of the complex pattern of the leaf is self-controlled, as shown by the fact that isolated leaf primordia in culture appear to go through all the normal stages of development, even though the resulting leaves are minute.

LEAF SHAPE

The very wide range of variation in leaf shape in seed plants needs no emphasis. The shape of the mature leaf is determined by three factors: (i) the shape of its primordium; (ii) the number, distribution and orientation of cell divisions; (iii) the amount and distribution of cell enlargement.

The form of the early leaf primordium varies considerably from species to species. As we have seen, in tobacco the young primordium has a simple, finger-like structure, but in maples (*Acer*), the development of the primordium forming the mid-rib is shortly followed by the appearance of two lateral branches at its base, and these three finger-like structures give rise to the main veins of the leaf. In a compound leaf such as that of ash (*Fraxinus*), a number of lateral lobes are formed from the central primordium, and these give rise to the leaflets of the mature leaf. The subsequent development of each leaflet resembles that of a simple leaf.

The comparative growth rates of the lamina and of the main veins have a profound effect on the ultimate form of the leaf. If lamina growth keeps pace with that of the main veins, then a leaf of simple outline results. On the other hand, if growth is more vigorous near the veins than in the other regions, then a lobed leaf will be formed, the ultimate shape depending also upon the pattern of vein development. In maple, for example, the early localized lamina growth around the main veins is normally followed by a wing-like growth to form a continuous sheet of lamina joining the veins, so that a *palmate* leaf is produced, but in some genetical variants the growth of the lamina is restricted more to the regions adjacent to the veins, so that a more "dissected" type of leaf is formed.

Leaf shape may be profoundly modified by environmental factors; for example, the submerged leaves of some aquatic plants, such as *Sagittaria* and *Ranunculus* spp., have a very different form from that of the aerial leaves. In certain species a considerable number of genes which cause wide variations in leaf shape are known. The successive leaves formed on

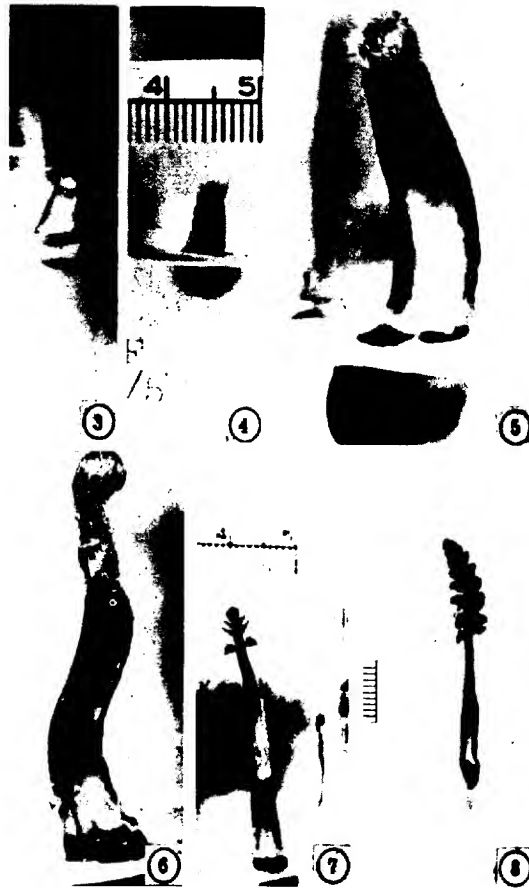


FIG. 2.12. Sterile culture of isolated leaf primordia of the fern, *Osmunda cinnamomea*, on a simple medium. Successive stages in the development of the leaf, from the earliest stage (top left). (From J. D. Caponetti and T. A. Steeves, *Can. J. Bot.* 41, 545, 1963.)

the stem from the seedling stage onwards very commonly show characteristic changes in shape (p. 246).

DIFFERENTIATION IN THE STEM

As we have seen, the cells in the apical meristem itself are generally small, densely cytoplasmic, have large nuclei, and are non-vacuolated. As we pass downwards from the apex to the regions in which cell vacuolation and differentiation begins to be apparent in the pith and cortex, we notice that there is a zone of cells between these two latter regions which is characterized by smaller, deeply staining cells, as seen in cross-section (Fig. 2.13).

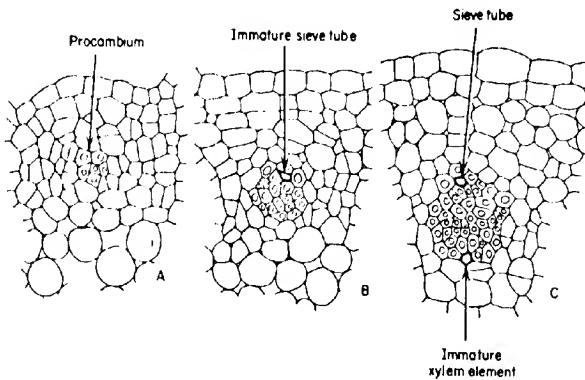


FIG. 2.13. Successive stages in the development of the procambium (cells with nuclei) in transverse sections of a stem of *Linum perenne*. (All $\times 430$. From K. Esau, *Amer. J. Bot.* **29**, 738, 1942.)

This latter zone gives rise to the future procambium. The level at which the procambium can be recognized varies considerably from species to species, but it commonly can be recognized in the zone of leaf initiation. At the highest level these densely staining smaller cells may be seen to form a complete ring, but not all the cells of the ring are destined to form the future vascular tissues and lower down the stem it can be seen that the ring has become broken into discrete strands, by vacuolation of some of the intervening cells of the former ring. These strands constitute the first clearly delimited procambium. Although the cells of the strands appear relatively small in transverse section, in longitudinal section they can be seen to be elongated and spindle-shaped.

As we have seen (p. 34), the development of the procambium is closely associated with leaf development. From the leaf base the procambium develops both acropetally into the leaf and basipetally to connect up with other vascular strands in the stem (Fig. 2.14).

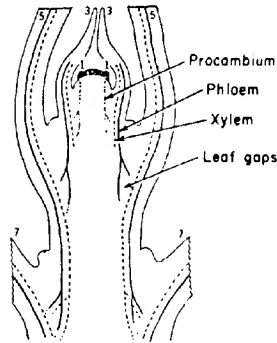


FIG. 2.14. Diagram illustrating the initial vascular differentiation in a shoot with a decussate leaf arrangement, as seen in longitudinal section. Continuous lines indicate phloem and broken lines indicate xylem. (From K. Esau, *Plant Anatomy*, John Wiley & Sons, Inc., New York, 1953.)

THE SHOOT APEX AS A SELF-DETERMINING REGION

The observation that procambial strands develop acropetally into leaf primordia at a very early stage of their development raises the question as to whether the procambium plays a role in determining the initial siting of leaf primordia. If this were the case, then it would imply that the activities and organization of the shoot apex are influenced and controlled by the already differentiated regions of the shoot. However, several lines of evidence seem to argue against this conclusion.

Firstly, it is clear that an organized shoot apex must arise *de novo* during the development of the embryo, and this may occur in free cell cultures in which the developing embryo is independent of the influence of any surrounding vascular tissues (p. 149). Similarly, shoot buds may arise spontaneously from undifferentiated tissue; for example, if chicory (*Cichorium*) or dandelion (*Taraxacum*) roots are cut in pieces, under appropriate conditions they will regenerate shoot buds from the cut surface at the upper end (Fig. 0.00). Adventitious buds will also develop in callus cultures of various species (p. 148). These observations suggest that the shoot apex represents a stable configuration of cells which will as it were, "crystallize out" from an undifferentiated mass of tissue under appropriate conditions. Thus, the shoot apex appears to be a self-organizing region.

Further evidence that the shoot apex is a self-determining region, and that it behaves as an organized whole, is provided by the observation that when the apex of *Lupinus alba* is divided into four sectors by vertical radial cuts through the centre, each of these sectors regenerates into a normal apex.

Other experiments have included the surgical isolation of the apex from surrounding tissues by four vertical incisions, so that it stands on a plug of parenchymatous tissue (Fig. 2.15). Under these conditions, in which the influence of the vascular tissue of the older parts of the shoot was removed, the apices of both *Lupinus* and the fern, *Dryopteris*, continued to

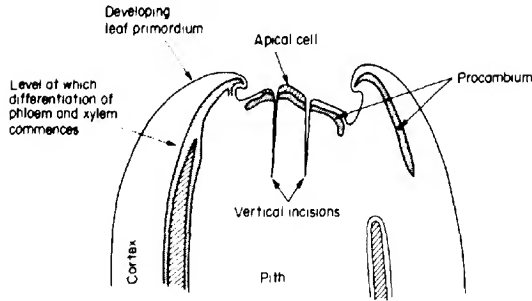


FIG. 2.15. Experiment involving surgical isolation of the shoot apical meristem of *Dryopteris aridata* from existing vascular tissues. Longitudinal section showing two of the vertical incisions which isolated the apical meristem on a plug of parenchymatous pith cells. Consequently, the only connection left between the apical meristem and existing vascular tissues was via parenchyma, yet the apex continued to grow, produced new leaf primordia, and developed normal vascular tissues. (Redrawn from C. W. Wardlaw, 1947.)

behave in a perfectly normal manner and produced new leaf primordia in the normal phyllotactic sequence, indicating, once again, the self-determining properties of the shoot apex. Indeed, so far from being controlled by the acropetal development of procambial tissue, there is much evidence that developing buds and leaves exert a stimulatory effect on the differentiation of vascular strands in the stem tissue below. For example, if young leaf primordia are removed from a shoot apex at a very young stage, vascular tissue fails to develop in the stem or, in the case of ferns, may be greatly modified or reduced. On the other hand, in callus cultures of chicory or lilac there is no differentiated vascular tissue, but if a bud is grafted into the callus, then vascular strands develop in the callus below the bud (p. 118).

The stimulatory effect of a bud on the development of vascular tissue appears to be due to the hormones, especially auxins and gibberellins, which it produces (p. 118). There is also evidence that hormones play an important role in the regeneration of vascular tissue (p. 118).

From these various types of evidence it is apparent that in respect of many of its activities the vegetative shoot apex is a self-determining region. On the other hand, when flowers are initiated, so that a vegetative shoot apex is converted into a flowering one, it appears that in many species this transition occurs under the influence of a stimulus arising within the mature leaves in the older parts of the plant (p. 205).

ROOT APICES

The apical region of roots shows both similarities and differences in comparison with shoot apices. No lateral organs such as leaves are initiated at the root apex, and hence growth is more uniform and there is no division into nodes and internodes. On the other hand, the

apex of the root is covered by a root cap, which is not represented in the shoot apex.

It is possible, by careful study of the patterns of division in the root apical region, to trace back the origin of the main zones of the differentiated root, the epidermis, cortex and vascular cylinder, to certain groups of initial cells in the main zone of cell division, the *promeristem* (Fig. 2.16A). It has proved remarkably difficult to identify the initial cells with certainty, however, and there is still considerable difference of opinion as to the number of initial cells in roots. Some authors have claimed that there are relatively few initial cells—perhaps only three, or even one. On the other hand, Clowes has produced evidence that, in the root tip of *Zea mays* and other species, there is a group of rather inert cells which constitute the *quiescent centre* (Fig. 2.16B) and that the actively dividing initial cells occur at the boundary or “surface” of the quiescent centre. The zone of actively dividing cells takes the form of an inverted “cup”. Several techniques have been used to study this problem, including autoradiographic studies to show the zones of active DNA synthesis and cell division, using a radioactive DNA precursor, such as ^3H -thymidine. Nuclei which show active DNA synthesis will incorporate the ^3H -thymidine, whereas inert cells will show no incorporation (Fig. 2.17). In this way, the existence of the quiescent centre can be clearly shown in various species. The function of the quiescent centre is still obscure.

The initial cells and their immediate daughter cells are non-vacuolated and cell division proceeds actively in this zone. Further back along the root, however, cell division becomes less frequent and cell vacuolation and extension commence. In the roots of many species (e.g. wheat) growth is fairly sharply separated into regions of cell division and cell extension, but in others (e.g. beech, *Fagus sylvatica*) there is a certain amount of division in cells which are beginning to vacuolate.

The boundaries of the future vascular cylinder, cortex and epidermis of the root become recognizable at a short distance back from the apical initials. These zones can be distinguished by the sizes of the cells and by the planes of division; the cells of the inner layers of the cortex tend to develop by periclinal divisions, whereas the planes of the cell walls of the future vascular tissue are less regular.

The phloem procambium becomes recognizable at an early stage, by virtue of its small cells as seen in transverse section. The procambium develops acropetally and the differentiation of xylem and phloem follows in the same direction, the phloem preceding the xylem (Fig. 1.1).

It was shown earlier that, in many respects, the shoot apex is a self-determining region, and this appears to be true also for the root apex. Thus, the pattern of vascular differentiation appears to be controlled by the apex itself. For example, if the apical 2 mm of roots is cut off, and the tip is turned about its longitudinal axis and replaced on the stump, the vascular tissue which later differentiates in the tip is out of line with that of the stump. Again, Torrey cut off the extreme tips of roots of *Pisum sativum* and grew them in a suitable culture medium. It was found that whereas the original roots showed triarch xylem (i.e. it showed three protoxylem groups), a certain proportion of the regenerated roots showed a diarch structure. Thus, the experimental treatment destroyed the original pattern of differentiation and yet a new one was determined by the meristem.

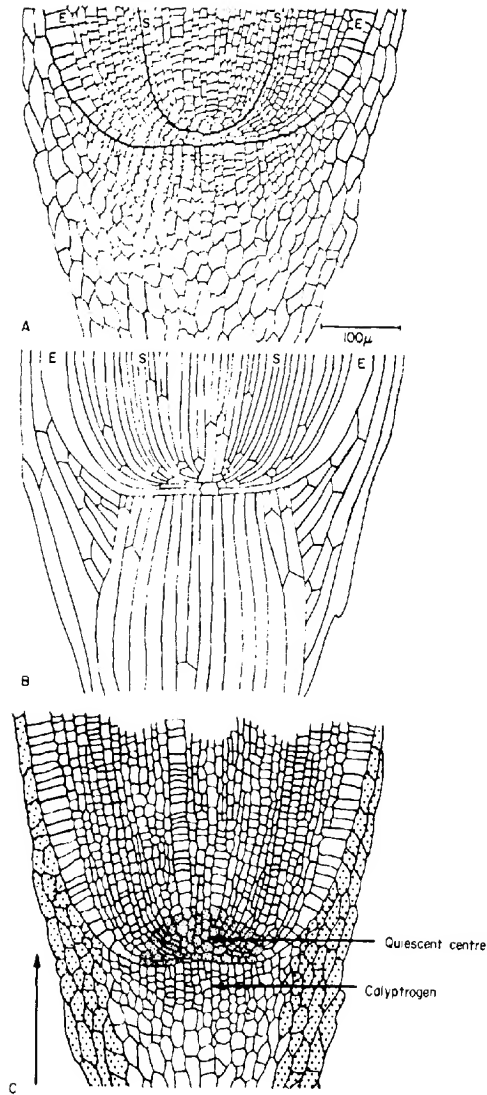


FIG. 2.16. A, B. Median longitudinal section of the root of maize (*Zea mays*). In A the outlines of the individual cells are shown, while in B the cell-lineages which can be traced back to the meristem region are outlined. E, epidermis; S, outer layer of stele. (From Clowes, 1961.) C. Longitudinal section of the root of maize (*Zea mays*), showing the position of the quiescent zone. (From Clowes and Juniper, 1968—see Further Reading.)



FIG. 2.17. Autoradiograph of root tip section of *Sinapis* photographed by dark ground illumination. The silver grains (white dots) are clustered over the nuclei that have synthesized DNA during the 48 hours in which tritiated thymidine was supplied to the plant. Note the quiescent centre in which no nuclei have been labelled. (From Clowes and Juniper, 1968.)

FLOWER INITIATION AND DEVELOPMENT

Sooner or later, one or more of the vegetative apices of a plant cease to produce leaves and buds and become converted into flowering apices. This transition involves a basic change in the structure of the shoot apex. The first detectable change is an increase in cell division in the region of the corpus between the central mother zone and the rib meristem (Fig. 2.18A), and this increased cell division gradually spreads to the central mother

zone and downwards into the flank regions. At the same time there is a marked reduction in cell division and growth in the rib meristem and pith where some of the cells undergo vacuolation. As a result of these changes, the shoot apical region is transformed into a structure consisting of a central pith of vacuolated cells covered by a "mantle" of smaller, densely staining meristematic cells (Fig. 2.18). During these changes the height of the apical

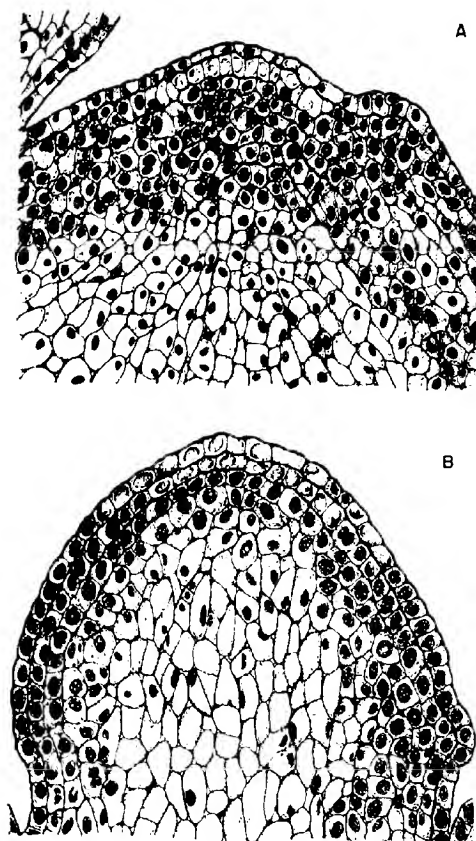


FIG. 2.18. A. Median section through the shoot apical meristem of *Xanthium strumarium*, during the transition from the vegetative to the reproductive state. Active division has commenced in the region between the central mother zone and the rib meristem and the zonation characteristic of the vegetative apex (Fig. 2.5) has been partly obliterated. B. Early reproductive apex of *Xanthium*, showing the meristematic "mantle" overlying the enlarged central rib meristem. (From F. B. Salisbury, *The Flowering Process*, Pergamon Press, Oxford, 1963.)

region increases considerably in most species, but where the inflorescence is a capitulum, as in the Compositae, it may become flattened.

The mantle, which includes both the tunica and the outer layers of the corpus, gives rise to the bracts and the flower primordia. Ultimately the flower primordia extend over the whole surface of the apex, so that all the meristematic tissue becomes differentiated. Thus, the structure of a vegetative apex becomes obliterated by the transition to a reproductive apex, and we have a change from an apex capable of unlimited growth to the determinate meristem of the inflorescence.

The subsequent pattern of development of the individual flower varies considerably from species to species. According to the "classical" viewpoint, the receptacle of the flower is a modified vegetative shoot and differs from the latter in that it is no longer capable of unlimited growth and has very short "internodes". The development of a relatively "primitive" flower, such as that of a buttercup (*Ranunculus*) (Fig. 2.19), bears out this view, since we find that during the early stages the apex still retains essentially the same structure as that of the vegetative shoot. Moreover, the initiation and early development of the various parts (perianth, stamens, carpels) are very similar to those of leaves, although the patterns of development diverge later. The stamen arises as a small protuberance and as it enlarges it gradually assumes the four-lobed form of the mature anther. The filament arises late in development, as the flower opens.

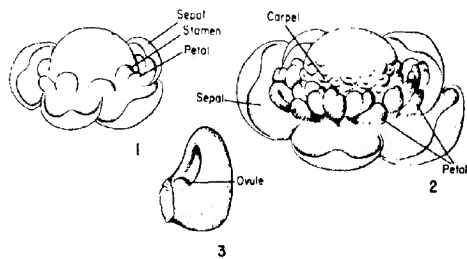


FIG. 2.19. Development of the flower in *Ranunculus trilobus*. 1 and 2. Two stages in the development of the entire flower. 3. Developing carpel. (Adapted from Payer, *Traité d'organogénie comparée de la fleur*, Paris, 1857. Reprinted from A. Fahn, *Plant Anatomy*, Pergamon Press, 1967.)

In flowers with an apocarpous gynoecium, i.e. with free carpels, the first stage of carpel development is the appearance of a rounded primordium similar to that of the other organs. This primordium elongates and a depression appears in the tip. As a result of further unequal growth each carpel adopts a horseshoe form (Fig. 2.19); these structures grow upwards and their margins meet and fuse. In flowers with a syncarpous gynoecium, the carpels may arise independently at first and fuse later, or they may be already joined from the earliest stages.

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CHAPTER 3

Plant Growth Hormones and Their Metabolism

THE GROWTH of a plant is a dynamic and complex, yet strictly controlled, process. This means that growth in different parts of the plant must be integrated and co-ordinated and we shall meet a considerable number of examples of such *growth correlations* in later chapters. The co-ordination of growth between different parts of the plant must clearly involve some control mechanism. Moreover, we have seen, in earlier chapters, that the development of organs, such as leaves or stems, involves an orderly sequence of phases of cell division and cell extension, so that there is also co-ordination of growth in *time*. As a result of intensive studies extending over many years, it is now known that hormones play a vital role in the control of growth, not only within the plant as a whole, but apparently also within individual organs. It is now realized that there are at least three major classes of growth-promoting hormones—*auxins*, *gibberellins* and *cytokinins*. In addition, other classes of plant hormones exist, particularly the “growth inhibitors” such as *abscisic acid* (ABA), but also including a gas, *ethylene*, which is apparently involved in many growth phenomena.

Growth hormones are translocated within the plant, and influence the growth and differentiation of the tissues and organs with which they come into contact. This leads us to a consideration of the nature and role of growth hormones. The word “hormone” was first used by animal physiologists, to refer to a substance which is synthesized in a particular secretory gland and which is transferred in the blood or lymph to another part of the body where extremely small amounts of it influence a specific physiological process. However, plant hormones differ in certain respects from the classical concept of hormones which was originally based upon the discovery of these substances in animals. In an animal a hormone is a substance produced in one particular organ such as a gland, and which is secreted to produce its typical and usually specific effect at a site distant from its point of origin. In the case of plant hormones we cannot always differentiate so clearly between the site of hormone synthesis and its place of action, although there is much evidence, referred to in Chapters 5 and 7, that plant hormones do usually have effects at sites distant from their place of production. Another difference between animal and plant hormones is that whereas the effects of most animal hormones are rather specific, a plant hormone can elicit a wide

range of responses depending upon the type of organ or tissue in which it is acting. For reasons such as these, plant growth hormones have frequently been referred to by other names, such as "growth regulators" or "growth substances". In general, though, the term "growth hormone" appears to be more appropriate, despite the difficulties discussed above, since it does intimate that these substances are active in extremely small quantities, and that in many instances they exert control over processes in tissues different from those in which they are synthesized.

Each of the chemically different categories of growth hormone has characteristic influences on growth and differentiation in plant cells and tissues. Auxins were the first plant growth hormones to be discovered, and consequently we shall now review very briefly the history of the discovery of auxins and their chemistry, followed by similar considerations of gibberellins, cytokinins, ethylene, and finally abscisic acid.

AUXINS

The basis of our modern knowledge of auxins lies in the work of Charles Darwin, published almost a century ago in a book entitled *The Power of Movement in Plants*. Darwin investigated the phenomenon of *phototropism*, the bending of plant organs in response to unilateral illumination.

Darwin experimented with seedlings of the ornamental canary grass (*Phalaris canariensis*). The coleoptile of grass seedlings proved a very convenient subject for the study of phototropism by Darwin and many other later workers. However, it was Darwin who first realized that the tip of the coleoptile *perceives* the unilateral light stimulus, but that the curvature *response* occurs lower down (Fig. 3.1). Darwin concluded that, "when seedlings are freely exposed to a lateral light some influence is transmitted from the upper to the lower part, causing the latter to bend". It was left to later researchers following Darwin to find out the nature of the "influence".

Various workers, particularly Boysen-Jensen and Paal, conducted experiments in the second two decades of this century which demonstrated that the growth-promoting influence transmitted from a coleoptile tip was of a purely chemical nature (Fig. 3.1). Paal was led to suggest that this chemical, under conditions of darkness or uniform illumination, continually moves down the coleoptile on all sides and acts as a *correlative growth promoter*.

The first successful isolation of the chemical messenger from coleoptile tips was carried out in 1926 by a Dutchman, F. W. Went, who thus extended the work of Boysen-Jensen and Paal. Went found that if he placed an excised oat (*Avena*) coleoptile tip upon a small block of agar gel, then the agar block acquired growth-promoting properties, in that if the block was separated from the tip and placed on one side of a coleoptile stump, then curvature of the stump resulted. An agar block which had not been in contact with a tip had no such effect. The conclusion was, therefore, that the chemical messenger had diffused from the tip into the agar block. Subsequent placing of the agar block on to a coleoptile stump allowed diffusion of the messenger out of the block and into the stump. However, this was

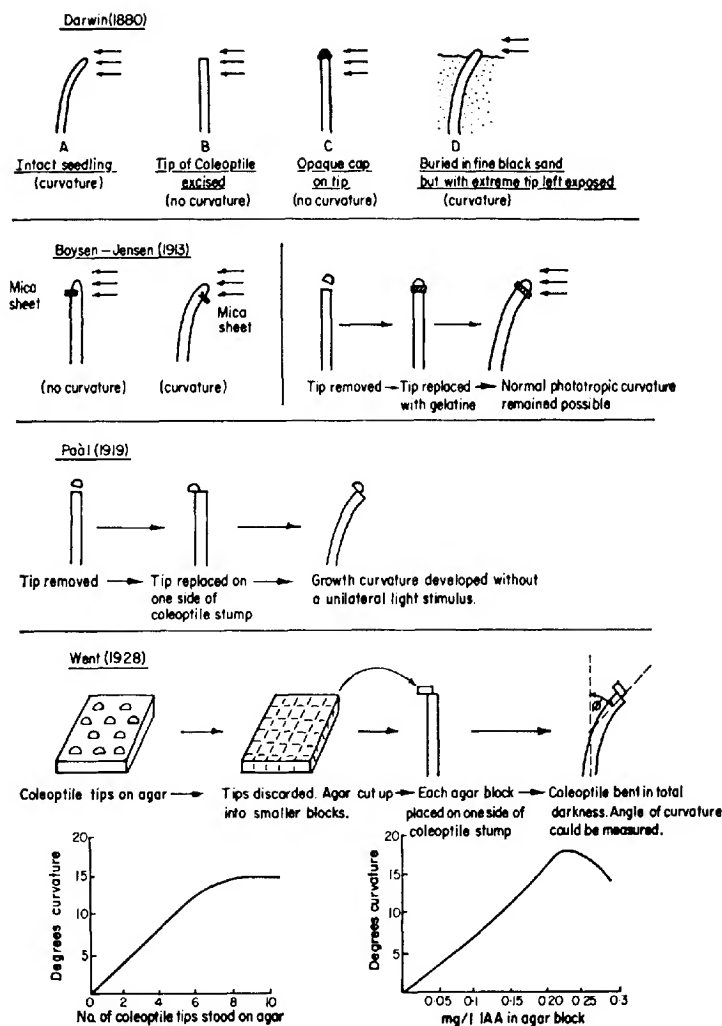


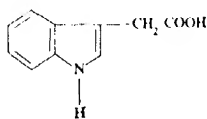
Fig. 3.1. A summary of important experiments which established the existence of auxin in plants. All experiments performed with coleoptiles of grass seedlings. Triple arrows indicate the direction of incident unilateral light. In the case of Went's experiments, dose-response curves are shown for the Went *Avena* curvature test; one for response against number of coleoptile tips diffused on agar, and the other for response against concentration of indole-3-acetic acid (IAA) in the agar block.

not only the first separation of a growth hormone from a plant but it also afforded Went a technique whereby he was able to make quantitative measurement of the growth hormone. The method of measurement that Went devised is a biological assay (*bioassay*) based on the curvature of a coleoptile stump in response to an asymmetric placing of an agar block containing the growth hormone (Fig. 3.1).

The name *auxin* (from the Greek *auxon*, to grow) was given to the growth hormone produced by the tip of a coleoptile. However, it is now known that auxins are present in *all* higher plants and not only in grass seedlings. Auxins are, as we shall see later, of fundamental importance in the physiology of growth and differentiation. They appear to be synthesized mainly in meristematic tissues such as those of the stem and root apex, young developing leaves, flowers and fruits.

The Isolation and Chemical Characterization of Auxin

Early attempts to isolate and chemically identify auxin suffered from the difficulty that it is present in plant tissues in extremely small quantities. It was found to be impossible at that time to obtain sufficient auxin from plants to prepare pure crystalline auxin for analysis. In fact the first crystalline auxin was obtained from human urine. Some confusion arose as a result of early attempts to determine the chemical nature of this auxin, but in 1934 it was shown to be indole-3-acetic acid (often abbreviated to IAA).



Indole-3-acetic acid (IAA)

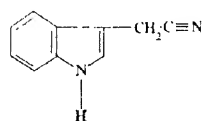
Since the initial discovery of IAA as an auxin, it has been found that this substance occurs in most plant species, and it is now believed to be the principal auxin in higher plants.

Much research is still conducted with the aim of elucidating the true nature of endogenous (i.e. internally synthesized) auxin in plants, and this involves the use of a number of modern chemical and physical techniques, as well as biological methods. As we saw earlier, the first isolation of auxin from plants was achieved by allowing diffusion of the hormone to take place from the plant tissue into a suitable inert medium such as agar gel. This method is still utilized and auxin obtained in this way is referred to as *diffusible auxin*. More commonly, plant tissues are extracted by organic solvents, such as di-ethyl ether or methanol. There are often quantitative and qualitative differences between auxins obtained by diffusion and extraction, even from the same tissue.

Chromatographic separation of the constituents of a plant extract results in the original extract being split up into a number of separate fractions. Each of these fractions is then

examined, to see whether or not it contains any auxin. To do this some sort of bioassay is usually used, such as the Went *Avena* test described earlier. There are numerous other types of biological assay for auxin that have been developed over the years, all of them making use of some measurable aspect of the effects of auxin in plants. One of the most commonly used tests is the coleoptile straight-growth test, in which sections of uniform length (say 5 mm) are cut from young *Avena* coleoptiles and placed in the extract to be tested for growth-promoting activity. However, although biological assays are indispensable in such studies, chemical identifications of the auxin must eventually be carried out. Even today this identification presents great difficulties, mainly because of the difficulty in obtaining sufficient auxin for analysis. Nevertheless, chemical characterization of auxins in plants is now possible, largely through the development and application of sensitive physico-chemical techniques such as ultra-violet and infra-red spectroscopy, and more recently mass-spectrometry has been applied to the problem.

The results of such studies have, in general, confirmed that IAA occurs as an auxin in plants, although it has become apparent that there are other substances in plants which also possess auxin activity. In many cases these other substances are indoles, closely related chemically to IAA; for example, the compound indole-3-acetonitrile (IAN) is known to be present in a number of plant species, particularly members of the Cruciferae, and pea plants appear to contain 4-chloro-indole-3-acetic acid as a natural auxin. There are also



Indole-3-acetonitrile (IAN)

compounds, such as phenylacetic acid, present in plants which are not indoles and yet possess auxin activity, but both the chemical natures and the physiological significance of most of these non-indole endogenous auxins remain obscure at the present time. It is because of the existence of more than one type of chemical which show auxin activity, that we often speak of auxins, rather than the singular, auxin.

Indolic Auxin Metabolism

Investigations of the metabolism of endogenous auxin have concentrated on the origin and fate of IAA in plant tissues. The reasons for this are first, the belief that IAA is the principal auxin, and secondly the concern to understand normal plant growth regulatory mechanisms. Since it is known that the *concentration* of auxin available to tissues can have a determining effect on growth and differentiation, then the factors which limit the concentration of IAA in plant tissues have received most attention. These factors include:

1. IAA synthesis.
2. IAA destruction.
3. IAA inactivation by processes other than destruction of the molecule.
4. Regulation of IAA transport between tissues and organs (see Chapter 5).

Auxin biosynthesis. As soon as IAA was identified as an endogenous auxin, it was suggested that it is formed from the amino acid tryptophan, a compound with an indole nucleus which is universally present in plant tissues, both in the free state and incorporated into protein. Indeed, it has been demonstrated many times that higher plants, or enzyme preparations from them, are able to bring about the conversion of exogenous tryptophan to IAA. Greater amounts of IAA are formed from added tryptophan in non-sterile plants or enzyme preparations than under sterile conditions, and in the late 1960s it was suggested that all the IAA present in plants is synthesized by epiphytic bacteria. Subsequent research has, however, revealed that completely sterile plant tissues and enzyme preparations are able to convert tryptophan to IAA. There seems little doubt, therefore, that IAA is synthesized by higher plants, utilizing a biosynthetic pathway involving indole-3-pyruvic acid (IPyA) and indole-3-acetaldehyde (IAAld) as intermediates (Fig. 3.2). In certain species, such as oat, tobacco, tomato and barley, there is some evidence that IAA can also be synthesized from tryptophan via tryptamine and IAAld (Fig. 3.2), but other species (e.g. pea, bean, cabbage, squash) do not contain tryptamine. A third pathway for IAA synthesis exists in members of the Cruciferae, in which tryptamine may be converted to indoleacetaldoxime and thence to IAN either directly or via the thioglucoside, glucobrassicin, followed by the conversion of IAN to IAA by the enzyme nitrilase. The relative importance of these alternative pathways for IAA synthesis is not known.

Apart from the indole compounds shown in the schematic representation of IAA synthesis in Fig. 3.2, a number of other indoles are known to occur naturally in plants. It is possible that any one or all of these indoles could serve as precursors of IAA, but clearly we still lack good information on the biosynthetic pathways for IAA actually occurring *in vivo*.

IAA destruction. It is well established that IAA is in one way or another fairly readily inactivated by most plant tissues. It appears, therefore, that the concentration of IAA in plants is regulated not only by its rate of synthesis, but also by inactivation mechanisms. The indications are that IAA catabolism as well as biosynthesis may follow more than one pathway.

(a) *Photo-oxidation.* IAA in aqueous solution will soon decompose if left exposed to light. This photo-oxidation of IAA is greatly accelerated by the presence of many natural and synthetic pigments. Thus, it is possible that plant pigments absorb light energy which energizes the oxidation of IAA *in vivo*. If this is so, then the most likely pigments involved are riboflavin and violaxanthin, for these are commonly present in plants in relatively large amounts and they absorb light in the blue regions of the spectrum which have been found to be the most active in inducing photo-oxidation of IAA.

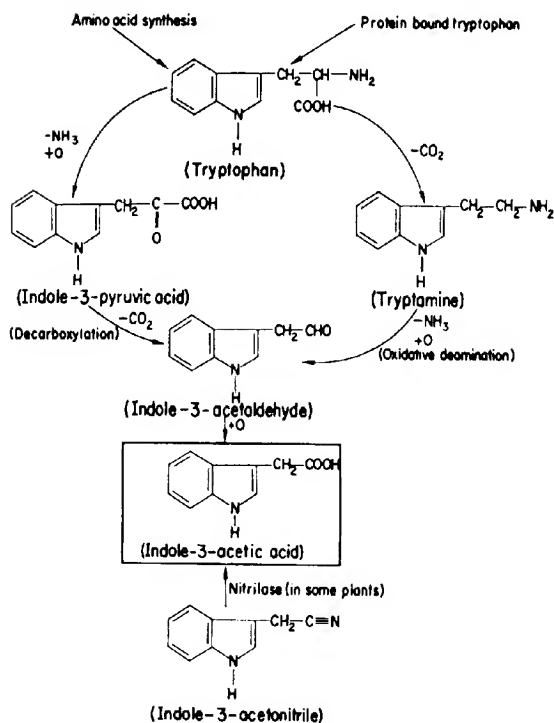
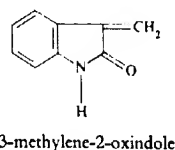
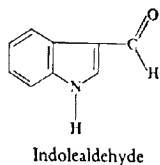


FIG. 3.2. Possible biosynthetic pathways for indole-3-acetic acid (IAA) in plants. (The pathway through tryptamine operates in some plant species, for example in oat, tobacco, tomato and barley, but not in others.)

Breakdown products of *in vitro* photo-oxidation of IAA include 3-methylene-2-oxindole and indolealdehyde, together with other unidentified compounds formed by cleavage of the indole ring.



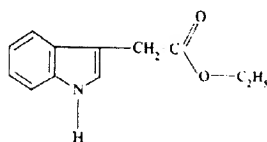
Nothing is known of the chemistry of IAA photo-oxidation within the plant, but indole-aldehyde and methylene-oxindole do occur naturally in many plants, and may perhaps represent a product of *in vivo* photo-oxidation. Some research has indicated that when methylene-oxindole is applied to plants, it may show growth-regulating properties, either inhibiting or accelerating the rate of growth. However, other more recent work has suggested that endogenous methylene-oxindole does not function as a growth regulator *in vivo*.

(b) *Enzymic oxidation of IAA.* Many plants contain an enzyme, or enzyme system, known as "IAA-oxidase", which catalyses the breakdown of IAA, with the release of CO_2 and consumption of O_2 . IAA-oxidase preparations from different plant species often have different properties, but they all show some similarities to the enzyme peroxidase. Full peroxidation of IAA (i.e. by H_2O_2 in the absence of O_2) does not occur, but oxygen is always required in addition to H_2O_2 . Addition of H_2O_2 to some IAA-oxidase preparations does enhance the rate of IAA degradation, but in other cases it does not; probably because crude plant enzyme preparations can possess the ability to generate H_2O_2 which would be available for peroxidase action. IAA-oxidase from higher plants requires manganese as a co-factor, and its activity is increased by monophenols and reduced by *ortho*- and *para*-dihydric phenols and polyphenols.

The pathway of IAA breakdown by IAA-oxidase is ill-understood. A principal product *in vitro* is 3-methylene-2-oxindole, and this is probably further metabolized to 3-methyl-2-oxindole. IAA-oxidase preparations that also contain cytochrome oxidase metabolize IAA to yield indole-3-aldehyde as the main product, and this latter compound is relatively stable compared with the oxindoles. The physiological significance of these observations is not at all clear at present. Furthermore, the relationships between IAA-oxidase and naturally occurring phenolics are still rather obscure, probably because many of the experimental studies have been made with relatively crude enzyme preparations. It is quite likely, nevertheless, that naturally occurring phenolic modifiers, or regulators, of IAA-oxidase activity do function in plants.

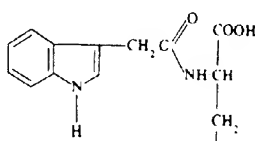
Interest in the enzymic oxidative destruction of IAA has been maintained by indications that such a process might be important in regulating auxin levels in plant tissues. Thus, there have been a number of reports that (1) IAA-oxidase activity rises with age of tissues, (2) there is a negative correlation between growth rate and IAA-oxidase content of various organs, and (3) that root tissues contain both very low IAA concentrations and very high IAA-oxidase activity. Nevertheless, there is not, as yet, conclusive proof that such correlations are physiologically important.

Inactivation of IAA by other Processes. Apart from degradation by photo-oxidation or enzyme activity, IAA can be inactivated in plant tissues by the formation of hormonally-inert complexes of various types. Thus, IAA is readily esterified by plant enzymes to yield its ethyl ester (indole ethyl acetate).



Indole ethyl acetate

Similarly, enzymic formation of conjugates such as indole acetyl-aspartic acid have been reported many times.



Indole acetyl-aspartic acid

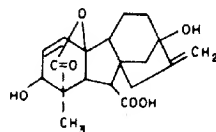
The nature of the esterifying or conjugating enzymes involved in these reactions remains very uncertain at present.

Conjugates are also formed between IAA and various sugars and sugar-alcohols, yielding compounds such as indoleacetyl-rabinose, indoleacetylglucose and IAA-myo-inositol, and between IAA and proteins.

GIBBERELLINS

The discovery of a group of plant growth-hormones now known as the gibberellins dates from the 1920s, when Kurosawa, a Japanese research worker, was investigating the "bakanae" ("foolish seedling") disease of rice caused by the fungus *Gibberella fujikuroi* (also known as *Fusarium moniliforme*). A characteristic symptom shown by rice plants infected by this fungus is an excessive elongation of stems and leaves, resulting in abnormally tall plants which usually fall over due to the spindly stem structure—hence, "foolish seedling". Kurosawa and fellow Japanese workers found that if they grew the fungus in a culture medium, and subsequently filtered off the fungus, then the culture filtrate, which was completely free of the fungus itself, would induce the same abnormal growth symptoms when applied to rice plants. It was clear that *Gibberella fujikuroi* secreted some substance into infected plants, or into the nutrient medium when grown in culture, which was stimulatory to stem and leaf elongation. In 1939 a small quantity of highly active crystalline material was isolated from such culture filtrates, and was given the name "gibberellin A". The chemical composition and structure of this material was not unequivocally worked out by the Japanese. It was not until 1954 that further progress was made, when British chemists

isolated and chemically characterized a pure compound from culture filtrates of *Gibberella fujikuroi*. They called this new substance *gibberellic acid* and found that it has the following structure:



Gibberellic acid when applied to many species of intact growing plants induced abnormally great extension of stems and leaves, but the response was found to be greatest when *genetic dwarfs* of various plant species were treated. Such treated dwarf plants assumed the appearance of the normal tall plants, from which the dwarfs had originally arisen by mutation (Fig. 3.3).

It might appear strange that a substance obtained from a fungus should produce essentially normal responses in higher plants. However, it is now known that gibberellic acid and substances very similar in both chemical structure and biological activity occur in healthy (i.e. non-infected) plants of all species. In fact, a number of these compounds have been isolated from higher plants, so that at the present state of knowledge there are over fifty chemically characterized compounds which produce effects similar to those elicited by gibberellic acid. These compounds are known collectively as the gibberellins, designated gibberellins A₁, A₂, A₃, A₄, etc. Gibberellic acid is numbered A₃. Some of the known gibberellins have been isolated from culture filtrates of *Gibberella fujikuroi*, and others from

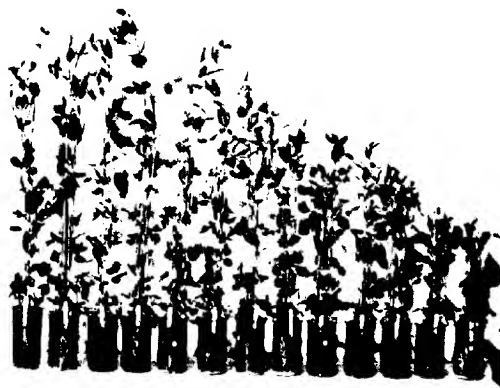
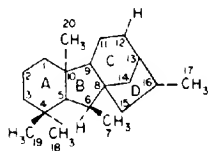


Fig. 3.3. Effect of gibberellic acid (GA₃) on shoot growth in dwarf pea plants (*Pisum sativum* c.v. Meteor). Plant at extreme right was not treated with GA₃, but the other plants received increasing doses of GA₃ from right to left. (Original print provided by Professor P. W. Brian.)

various organs of higher plants. All have the same basic molecular structure (the *ent*-gibberellane carbon skeleton) as gibberellic acid, differing from one another mainly in the number and positions of substituent groups on the ring system, and the degree of saturation in the "A" ring (see Fig. 4.5). It is highly likely that further additions will in future years be added to the list of gibberellins.



The *ent*-gibberellane carbon skeleton

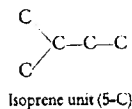
Apart from gibberellins having the gibberane carbon skeleton, there are some other compounds known which possess gibberellin-like biological activity but differ markedly in chemical structure. For example, a compound called helminthosporol isolated from the fungus *Helminthosporium sativum*, and phaeolic acid obtained from bean seeds.

The discovery that the gibberellins, or at least some of them, are natural growth hormones in higher plants, necessitated a complete reconsideration of views of the hormonal control of plant growth and differentiation. One could no longer think of development of cells and tissues as being influenced by only one growth hormone, that is auxin, but consideration had to be made of the effects of gibberellins and, later, other hormones (e.g. cytokinins) on the processes affected by auxins. Indeed, as we shall see later, auxins, gibberellins and cytokinins *interact* in their influences on plant growth and differentiation, and it is highly likely that abscisic acid and ethylene also interact with other growth hormones.

Gibberellin Metabolism

Gibberellins are chemically diterpenes, which are themselves members of a vast group of naturally occurring compounds in plants called terpenoids. Considerable knowledge of terpenoid biochemistry exists, and consequently rapid progress has been made in elucidating the outlines of gibberellin biosynthesis.

Gibberellin biosynthesis. All terpenoids are basically built up from "isoprene units", which are five-carbon (5-C) compounds. The linking together of two isoprene units yields a

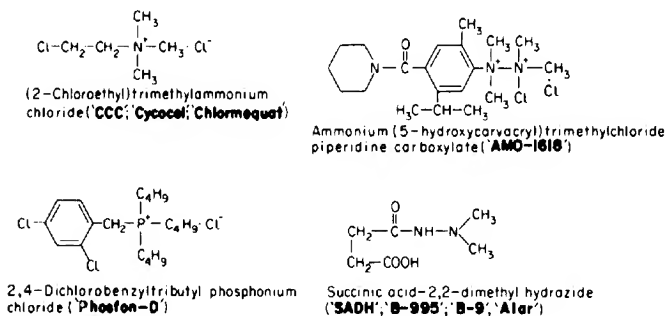


monoterpene (C-10), of three a sesquiterpene (C-15), of four a diterpene (C-20), and of six a triterpene (C-30).

A summary of the biosynthetic sequence of gibberellins in higher plants as we presently know it is shown in Fig. 3.4.

It is of interest to note here that the growth inhibitor abscisic acid is a sesquiterpenoid, and it is possible that the initial steps in the biosynthesis of gibberellins and of this substance may involve a common pathway from mevalonic acid.

A number of synthetic *growth retardants* have been discovered in recent years (e.g. below) :



Some of these are proving of considerable importance in agriculture (see p. 137). Several of these growth retardants have been shown to act by inhibiting gibberellin biosynthesis in the plants to which they are applied. For example, the growth retardant "Amo-1618" has been shown to inhibit the biosynthesis of gibberellin in homogenates of the endosperm of the wild cucumber (*Echinocystis macrocarpa*). It appears that Amo-1618 inhibits the cyclization of geranylgeranyl pyrophosphate to (–)-Kaurene (Fig. 3.4). The growth retardant "Cycocel", or "CCC" appears to act similarly.

Gibberellin catabolism. Very little is known of the eventual fate of gibberellins in plant tissues. There is evidence that gibberellins retain their physiological activity for some considerable time in plants. This is in marked contrast to the rapid inactivation of applied natural auxins (such as IAA) in plant tissues, but it is a common observation that even large doses of gibberellins are not particularly injurious to plants whereas auxins can be, perhaps due to an effect on the production of ethylene (p. 113). Thus, it would seem more important that plants should have the capacity for speedy inactivation of auxin when its concentration exceeds a certain value. However, there is evidence that considerable *interconversion* of gibberellins takes place in plant tissues. That is, one gibberellin can be converted into a different gibberellin (see Fig. 3.4). Moreover, there is evidence for the existence of gibberellin glycosides (i.e. conjugates with sugars) in plant tissues (e.g. 2-O-D glucopyranosyl-GA₃ is present in seeds and seedlings of *Phaseolus coccineus*), and these may represent the products of an inactivation mechanism.

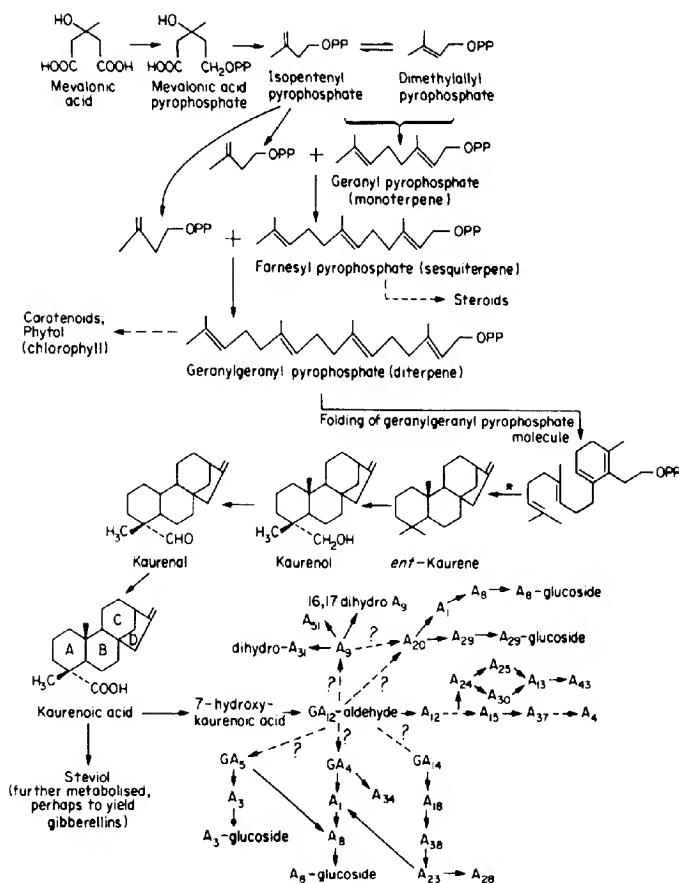


FIG. 3.4. Possible biosynthetic pathways for some of the known gibberellins from mevalonate. The sequence of reactions and transformations shown are those which appear to occur in higher plants; dashed lines indicate transformations which involve as yet unknown intermediates. In the fungus *Gibberella fujikuroi*, the pathways are similar but not identical to those in higher plants. The cyclization of geranylgeranyl pyrophosphate (*) is blocked by growth retardants such as CCC, Phosfon D and AMO-1618. (Diagram prepared with advice from Dr. J. Mac-Millan and Dr. Ian D. Raitlen.)

Gibberellic acid in solution can be decomposed by acid hydrolysis, particularly at higher temperatures, to yield compounds such as gibberellenic acid, allogibberic acid and gibberic acid. The latter compound does not retain any hormonal activity, but gibberellenic and allogibberic acids can still elicit some of the physiological effects of gibberellins.

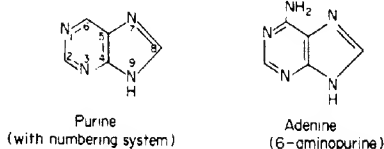
CYTOKININS

The discovery of this group of plant growth hormones came from work concerned with the *in vitro* culture of young plant embryos and tissue explants. Studies by Haberlandt in the first two decades of this century demonstrated the existence in plant tissues of a diffusible factor which stimulated parenchymatous cells in potato tubers to revert to a meristematic state. That is, cell division could be induced by the factor.

Many workers, particularly Skoog and Steward in the U.S.A., have made a close study of the growth requirements of callus-cultures (a mass of undifferentiated and usually rapidly dividing cells (p. 144)) prepared from the parenchymatous pith cells of tobacco, and from carrot roots. It is primarily as a result of their work during the 1950s that we became aware of the existence of cytokinins—plant growth hormones originally regarded as being particularly important in the processes of cell division and differentiation, but which more recently have been found to be implicated in various other physiological processes such as senescence and apical dominance.

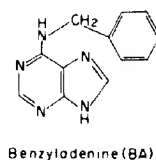
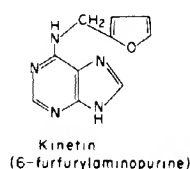
Skoog used a tissue culture technique in which an isolated piece of tobacco pith is placed on the surface of agar gel into which had been incorporated various nutritive substances and other, hormonal, factors. The exact composition of the agar medium was varied, and the effects on the growth and differentiation of the pith cells noted. For growth to occur it was found that it was necessary to add not only nutrients to the agar, but also hormonal substances such as auxin. However, when auxin (IAA) was applied alone with the nutrients very little growth of the pith explant occurred, and that which did consisted predominantly of cell enlargement; very few cell divisions occurred, and no differentiation of cells took place. If, however, the purine base *adenine* was incorporated into the agar medium along with IAA, then the parenchymatous cells were induced to divide and a large callus mass was created. Adenine added without auxin, however, did not cause cell division in the pith tissue. There was, therefore, an *interaction* between adenine and auxin, resulting in the triggering off of cell division. Adenine is a purine derivative (6-aminopurine) and is a naturally occurring component of nucleic acids.

Later, another substance with similar, but more potent, effects as adenine was prepared



from degraded deoxyribonucleic acid (DNA). This substance was found to be 6-(furfurylamino) purine, basically similar in structure, therefore, to adenine. Due to its property of actively promoting cell division (in conjunction with auxin) it was given the name of *kinetin*. Indole-3-acetic acid and kinetin were found to have interacting effects upon cell division and differentiation in tobacco pith cultures (Fig. 6.2), similar to those shown by IAA and adenine.

Kinetin is a synthetic cytokinin, which does not occur naturally in plants. A number of other synthetic cytokinins have been subsequently discovered, among the most active of which are the compounds benzyladenine (BA) and tetrahydropyranylbenzyladenine (PBA):



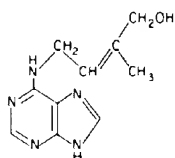
Endogenous Cytokinins

Although kinetin, BA and PBA have never been shown to be present normally in plants, substances which produce similar physiological and morphological effects have been found in various organs of many plant species, particularly in "nurse tissues" such as coconut milk (a liquid endosperm), in immature caryopses of *Zea mays*, and in immature fruits of *Aesculus hippocastanum* (horse chestnut), banana and apple. These naturally occurring substances, together with other synthetically prepared compounds which have effects on growth similar to those of kinetin, have been given the generic name of *cytokinins*. A cytokinin, therefore, is a substance which, in combination with auxin, stimulates cell division in plants and which interacts with auxin in determining the direction which differentiation of cells takes.

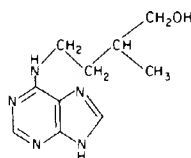
Most available evidence suggests that naturally occurring cytokinins are purine, particularly adenine, derivatives. In 1964 the New Zealander, Letham, isolated a cytokinin from

sweet corn kernels and identified it as 6-(4-hydroxy-3-methyl but-2-enyl) amino purine. For convenience, Letham has called this substance *Zeatin*.

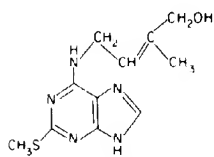
Since the first isolation and characterization of zeatin, a quite extensive range of naturally occurring cytokinins have been identified from various plant sources. All the known naturally occurring cytokinins are adenine derivatives (i.e. they are 6-substituted amino purines). Some of the naturally occurring cytokinins are illustrated below. Zeatin is the most active of the known natural cytokinins. The cytokinin (*o*-hydroxybenzyl) adenosine, isolated from poplar leaves, is of interest since the base is identical with that of the synthetic cytokinin benzyl adenine, apart from the additional hydroxyl group.



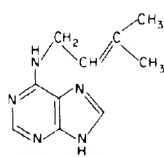
Zeatin



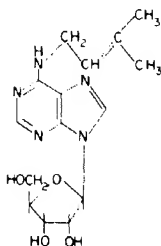
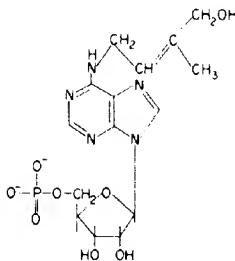
Dihydrozeatin



Methylthiozeatin

Dimethylallyladenine
(also known as isopentenyladenine)

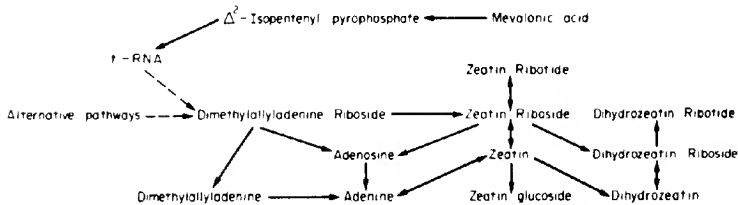
Like other purines, the natural cytokinins readily form the riboside (purine base + ribose sugar) and ribotide (base + ribose + phosphate group) derivatives, and may also, as we shall consider later, be contained in ribose nucleic acids. Examples of a naturally occurring cytokinin riboside and a cytokinin ribotide are shown below:

Dimethylallyladenine riboside
(=isopentenyladenosine, IPA)

Zeatin ribotide

(a) *Biosynthesis*. It may be assumed that the biosynthesis of natural cytokinins takes place by the substitution of characteristic side chains on to carbon 6 of the adenine moiety which is of common occurrence in plant cells. The side chain of a natural cytokinin contains five carbon atoms and this suggests that it is derived from the isoprenoid biosynthetic pathway. As we shall see later, cytokinin groups occur in certain species of t-RNA, and it has been shown that radioactivity from labelled mevalonate (MVA) becomes attached to specific adenine groups in the t-RNA, to give the dimethylallyl side chain of the cytokinin, IPA.

The occurrence of cytokinin groups in t-RNA means that free cytokinins may be formed from the degradation of t-RNA and this has, indeed, been shown to be the case. However, there is considerable doubt as to whether the observed levels of free cytokinins in actively growing plant tissues can be accounted for solely by degradation of t-RNA, and it may well be that there is another pathway of cytokinin biosynthesis in which the isoprenoid side chain is attached directly to free adenine. Experiments conducted to date, however, have failed to demonstrate incorporation of ^{14}C -MVA into the side chain of free cytokinins, and the significance of this is uncertain at present. Apart from direct substitution of adenine with an isoprene-derived side chain, at least some of the naturally occurring cytokinins are formed by conversion from one or other cytokinins. A possible mechanism of metabolic interconversion of natural cytokinins in plants is illustrated in Fig. 3.5.



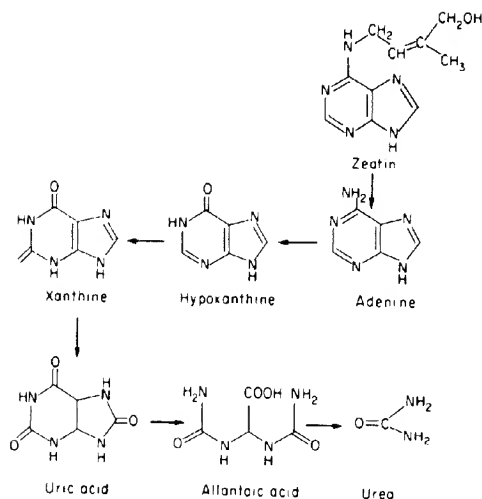
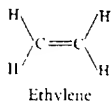


FIG. 3.6. Catabolism of the naturally occurring cytokinin zeatin. (Adapted from D. Schlee, H. Reinbothe and K. Mothes, *Z. Pflanzenphysiol.* **54**, 223-36, 1966.)

be a "storage" form of cytokinins from which the free bases may be released under certain conditions.

ETHYLENE

Ethylene may appear a curious substance to consider as a hormone. It is a very simple organic molecule, contrasting with the more chemically complex gibberellins, cytokinins,



auxins and abscisic acid. Moreover, ethylene exists as a gas at normal temperatures. Thus, ethylene, if a plant hormone, is a gaseous hormone. It can be, and has been, argued that there are theoretical advantages to the plant in having a gaseous, diffusible growth regulator in addition to other hormones which necessarily move through living cells to reach their site of action. Treatment of plants with very low concentrations of exogenous ethylene has many profound effects on their physiological and metabolic activities. However, evidence is also accumulating that endogenously synthesized ethylene is involved in the normal control of many aspects of plant growth, differentiation and responses to the environment.

It has been known for many years that developing fruits evolve ethylene, and that the time of maximum ethylene production by a ripening fruit coincides with the time of the *respiratory climacteric* (the latter term is applied to the large increase in respiration rate which occurs during the ripening period of many fruits, prior to a fall off in respiration as the fruit enters a senescent decline). It has been found that exposure of fruits to ethylene results in a hastened and enhanced respiratory climacteric with earlier ripening. In fact, ethylene is so effective in stimulating respiration that it does so even in fruits, such as oranges and lemons, which do not naturally experience a respiratory climacteric. The speeding of fruit ripening by ethylene has proved of great commercial value in the citrus industry.

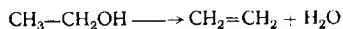
In recent years, however, it has become apparent that we must consider that ethylene has much wider physiological significance than its effects in fruit ripening alone would indicate. In general, it would appear that many effects previously considered to be induced directly by auxin may be mediated by an intervening step in which auxin leads to an increase in the formation of ethylene, following which ethylene induces the actual response.

In those situations where auxin-induced ethylene is responsible for effects of auxin, it is possible to regard ethylene as an intermediate hormone in much the same way that cyclic AMP serves as an intermediate in the action of many animal hormones. Not all auxin effects however, appear to involve ethylene as an intermediate. Neither can all effects of ethylene be elicited by auxins. Thus, ethylene cannot substitute for auxin in (a) the stimulation of cell elongation in the vast majority of plant species, (b) in promotion of growth in tissue cultures and, (c) in inhibition of senescence, ripening and abscission. Similarly, auxin usually cannot substitute for ethylene in the promotion of processes such as leaf senescence, abscission and seed germination.

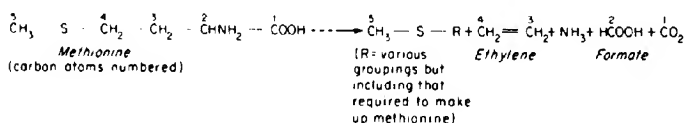
Ethylene can be accurately and sensitively measured relatively easily by use of gas-liquid chromatography, and it has been found that rates of ethylene biosynthesis in plant tissues vary widely from organ to organ and the stages of their development. In vegetative plants, the highest rates of ethylene production occur in most actively growing regions of stems and leaves, particularly in meristematic tissues. For example, in meristematic parts of the apical region of the stem of pea plants, ethylene production has been measured as $0.43 \mu\text{l hr}^{-1} \text{ kg}^{-1}$ fresh weight of tissue, whereas in older internodes only $0.04 \mu\text{l hr}^{-1}$ ethylene was produced per kg fresh weight. In general, parts of the plant rich in endogenous auxin also produce greatest quantities of ethylene, although senescent tissues usually produce relatively large amounts of ethylene and yet are low in auxin content (Chapters 5 and 12). Amongst the highest levels of ethylene production in plants are those recorded for ripening fruits of some species: passion fruits, for example, can produce $500 \mu\text{l kg}^{-1} \text{ hr}^{-1}$.

Ethylene Metabolism

Although ethylene synthesis in fungi appears to take place mainly by the dehydration of ethyl alcohol:

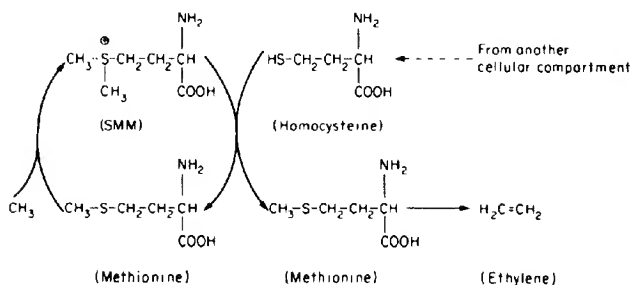


this pathway does not appear to operate in higher plants. Several substances have been suggested as precursors of ethylene in higher plants, but research over the past decade has, in the main, supported the view that the amino acid L-methionine is the principal, or perhaps sole, natural precursor. Addition of specifically radiolabelled methionine to plant tissues has shown that ethylene is formed principally from carbon atoms 3 and 4:



However, despite considerable work on the problem, it is not yet possible to put forward a definite scheme to illustrate intermediates in the biochemical conversion of methionine to ethylene in higher plants.

Recent studies by Hanson and Kende on ethylene production in senescing *Ipomoea tricolor* petals (see Chapter 12) have suggested the possibility that the rate of ethylene production in these tissues can be regulated by availability of homocysteine to a cellular compartment in which methionine is first formed from S-methylmethionine (SMM) and then is consumed in ethylene production according to the following scheme (this illustrates that SMM can act as a methyl ($-\text{CH}_3$) donor, and that during senescence of the petals the methyl group is transferred to homocysteine, to give two molecules of methionine. One of the two-product methionine molecules is remethylated to SMM, and the other contributes to a rise in free methionine level and hence to increased ethylene formation):



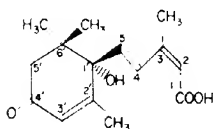
Until recently, it was not thought that higher plants possessed a mechanism for degrading ethylene. This did not seem incongruous because the ease with which the gas diffuses out of plant tissue (see p. 65) appears to provide a means for controlling endogenous concentrations. In other words, it seemed that the process of control of ethylene concentrations in

plants by emanation was analogous in effect to the process of degradation for other growth regulators. It has now been shown that in *Vicia faba* (broad bean) ethylene is metabolized very rapidly to ethylene oxide. Although the mechanism and generality of this process is not yet known its very demonstration has important consequences. Thus, most assessments of rates of ethylene biosynthesis have relied on the assumption that rates of emanation are proportional to rates of production (see above, p. 65). Clearly, at any rate in those plants which can metabolize ethylene, this assumption, and conclusions drawn from making it, are invalid.

ABSCISIC ACID

Studies on abscission and on dormancy in buds and seeds (Chapters 11 and 12) during the 1950s and early 1960s indicated the possible existence of a hormonal plant growth inhibitor. The chemical structure of the inhibitor present in fruits and leaves of cotton plants was eventually elucidated in 1965, and in the same year the same compound was isolated and identified from buds of dormant *Acer pseudoplatanus* trees.

The substance isolated from *Acer pseudoplatanus* and cotton was named abscisic acid (ABA), and proved to be a sesquiterpenoid (p. 57) of the following structure:



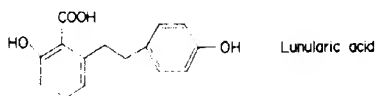
Absciscic acid (ABA)

(Revised absolute configuration. Numbering system for carbon atoms shown)

The molecule contains an asymmetric carbon atom (1') and therefore exhibits optical isomerism. However, only the (+) enantiomorph occurs naturally in plant tissues. Absciscic acid also shows geometric isomerism. Steric considerations demand that the side chain always be *trans* around carbon 5 of the side chain, but the molecule can be either *cis*- or *trans*- around carbon 2 of the side chain. Most of the ABA present in plant extracts is in fact (+)-2-*cis* ABA, though small amounts of (+)-2-*trans* ABA may also be present. By convention, the naturally occurring (+)-2-*cis* form is simply referred to as abscisic acid, or ABA.

Absciscic acid has been isolated from numbers of species of angiosperms, gymnosperms, ferns and mosses, but it does not appear to occur in liverworts. However, the compound *lunularic acid* has been identified in at least eight species of liverwort and some algae, and

may well play roles in the physiology of liverworts similar to those served by ABA in more highly evolved plants, for it is a very potent growth inhibitor and also appears to be involved with the mechanisms of dormancy and gemma growth in certain liverworts.



Absciscic Acid Metabolism

Because ABA is a sesquiterpenoid, it seems likely that the compound is synthesized in plants by operation of the common pathway for terpenoid biosynthesis as far as farnesyl pyrophosphate (Fig. 3.4). The feeding of plant tissues with radioactive mevalonate has been shown to result in the formation of radioactive ABA, and overall experimental evidence strongly suggests that ABA can be synthesized directly by the reactions of the isoprenoid biosynthetic pathway. However, it has also been found that certain naturally occurring carotenoids, such as violaxanthin, can be photo-oxidized to yield products that are very similar in structure to ABA (Fig. 3.7), and that these breakdown products may be further metabolised to ABA. Energization of the photo-oxidation of carotenoids to ABA-like compounds requires high light intensities, and for this and other reasons it is very unlikely

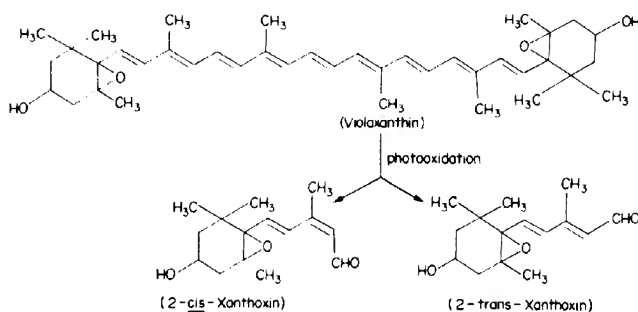
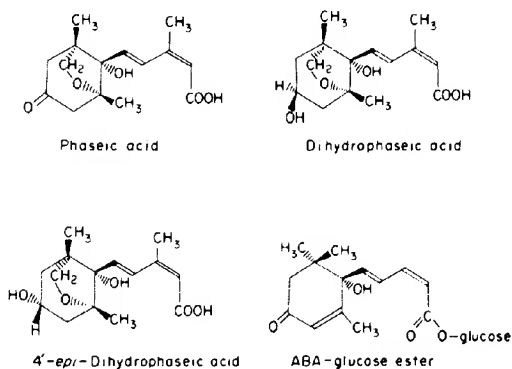


FIG. 3.7. Carotenoids such as violaxanthin can be broken down in the presence of strong light to yield products such as xanthoxin which are structurally similar to abscisic acid (ABA). It is possible that some of the ABA in plants arises by the further metabolism of carotenoid breakdown products, although most naturally occurring ABA is considered to be synthesized directly by the reactions of the terpenoid biosynthetic pathway.

that physiologically important ABA is formed in this way from carotenoids. For example, ABA occurs in etiolated plants and, furthermore, rapid rises in ABA levels occur in response to water-stress even in dim light or darkness. The principal means of ABA synthesis in plants therefore appears to be by direct synthesis from mevalonate via farnesyl pyrophosphate, although enzyme-mediated oxidation of violaxanthin to xanthoxin may also play some part in the biosynthesis of ABA in certain tissues.

The sites of synthesis of ABA within plants have not yet been investigated extensively, but indirect evidence suggests that most or perhaps all ABA is formed in mature green leaves and in fruits. ABA may be translocated from leaves to other regions such as the shoot apex and there inhibit growth and perhaps induce the formation of resting buds (see Chapter 11). There is also experimental evidence which indicates that plastids, particularly chloroplasts, may serve as centres of ABA synthesis.

As endogenous ABA levels fluctuate in relation to changes in growth rate, water potential and season, then not only synthesis but also inactivation of the molecule must take place in plant tissues. Relatively little is known of the factors concerned in the regulation of ABA inactivation, but it has been found that applied ^{14}C -ABA is rapidly conjugated in plants to form the glucose ester of ABA. This ester seems to be quite stable in plants, and possesses hormonal activity similar to that of ABA. Degradation of ABA also occurs, however, the early stages of which involve hydroxylation and oxidation of methyl substituents of the ring. Work conducted so far on the metabolism of applied 2- ^{14}C -(\pm)-ABA has indicated that in tomato exogenous ABA is rapidly converted into its glucose ester and phaseic acid, but in *Phaseolus vulgaris* the major metabolites have been identified as phaseic acid, dihydrophaseic acid, and 4'-*epi*-dihydrophaseic acid:



There is some evidence that phaseic acid may have regulatory functions in the physiology of plants, for example in the inhibition of photosynthesis in plants subjected to water stress.

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CHAPTER 4

Mechanism of Action of Plant Growth Hormones

ALTHOUGH almost a century of increasingly intensive research has passed since Darwin first recognized that a transmitted stimulus to growth is involved in the regulation of plant development, and approximately 50 years since a plant hormone, auxin, was first chemically characterized, we still do not understand the basic mechanism whereby plant hormones exert such large and varied physiological effects. Nevertheless, the voluminous accumulated information concerning the mechanism of action of plant growth hormones has served to formulate current views on the mode of action of these substances in molecular terms.

For various historical and practical experimental reasons, research into the mechanism of action of plant growth hormones has tended to concentrate upon particular physiological responses. As we consider below, for example, studies of the mechanism of auxin action have been largely concerned with its role as a regulator of cell extension growth, whereas equivalent research on gibberellins has concentrated on their effects on enzyme synthesis and secretion in aleurone cells of germinating cereal seeds.

We will now consider what is known of the mechanism of action of each of the categories of plant growth hormone, drawing appropriately upon information derived from the various experimental approaches that have been applied to the problem.

Relationships Between Molecular Structure and Hormonal Activity

The identification of indole-3-acetic acid (IAA) in 1934 as a naturally occurring plant growth hormone was followed by investigations to determine just what was "special" in the chemical structure of IAA to impart such profound influences on growth and developmental processes. It was hoped that an understanding of this would provide a lead into elucidating the mechanism of action of auxins within the cells. Similar considerations have more recently been applied to the gibberellins, cytokinins, ethylene and abscisic acid following their discovery.

Structure-activity relationships of auxins. The methods of studying this problem were initially largely empirical, in that many compounds were tested in suitable bioassays to find whether or not they possessed any "auxin activity". Some of these substances did in fact produce effects similar to those which IAA itself elicited, even when they were supplied at very low concentrations. Over the years a very large number of such compounds have been found, all of which have been synthesized in laboratories and are, therefore, called *synthetic auxins*. These synthetic auxins do not fall into any one particular class of compound, but despite the diversity of structure shown by the synthetic auxins, very serious efforts have been and still are being devoted to pinpointing exactly what attributes a molecule must possess for it to have activity as an auxin. It is hoped that elucidation of the molecular requirements for auxin activity will help in an understanding of the mechanism by which auxin operates in plant cells.

The first synthetic auxins found were compounds closely related to IAA (in having the indole ring) such as α -(indole-3)-propionic acid, α -(indole-3)-butyric acid and β -(indole-3)-pyruvic acid (Fig. 4.1) (which is now known also to occur naturally). However, many other synthetic auxins more markedly different in structure from IAA were subsequently discovered, and some of the more active of these, such as 2,4-dichlorophenoxyacetic acid (2,4-D), 2,4,5-trichlorophenoxyacetic acid (2,4,5-T) and 4-chloro-2-methylphenoxyacetic acid (MCPA) (Fig. 4.1), are not indole compounds.

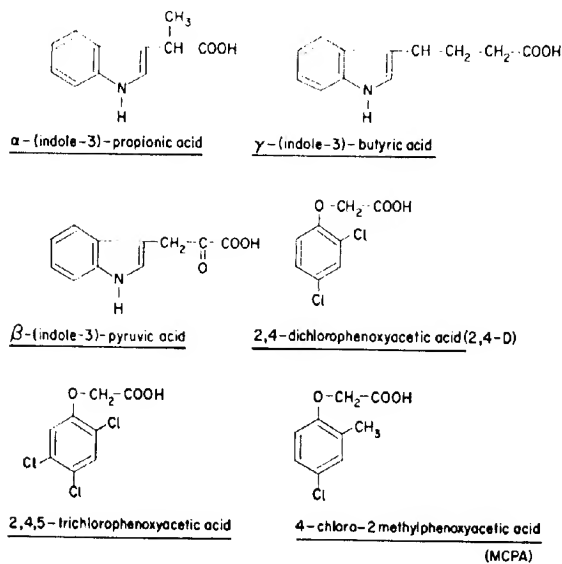


FIG. 4.1. The chemical configurations of some natural and synthetic auxins.

As a result of comparing the structures of all the then available synthetic and natural auxins, in 1938 a list was drawn up of the general structural requirements of a molecule for it to behave as an auxin. Thus, it was said that an active molecule must possess: (i) a ring system with at least one double bond present; (ii) a side chain containing a carboxyl group (or group easily converted into a carboxyl-group); (iii) at least one carbon atom between the ring and carboxyl group in the side chain; (iv) a particular spatial relationship between the ring system and the carboxyl group. Later on, it was thought that another requirement of molecules which have activity as auxins is that they must have the ability to form a covalent bond at a position on the ring system *ortho* to the side chain which terminates in a carboxyl group. An examination of the structures of the synthetic auxins shown in Fig. 4.1 will reveal that they all comply with these general requirements, but there are other compounds now known which possess auxin activity and yet do not fully comply with the above list of structural requirements. For example, certain benzoic acid derivatives are active auxins (Fig. 4.3), and yet have no side chain. On the other hand, the activity of certain thiocarbamates indicates that not even the unsaturated ring is essential, although it is necessary that these latter compounds should have a planar structure. Furthermore, instead of the ability to form a covalent bond at the *ortho* position, it is now known that the requirement is that there should be a fractional positive charge at a specific point on the ring (p. 75).

On the assumption that a carboxyl-terminated side chain, and a "free" *ortho* position on the ring system are essential for activity, it was proposed that the basic reaction of an auxin within the cell involves two parts of the molecule, the carboxyl group of the side chain and an *ortho* position of the ring system. This led to what is called the "two-point attachment theory" for auxin action. The research workers who put forward this theory proposed that there is covalent bond (i.e. chemical bond) formation at these two points between the auxin molecule and some constituent, possibly a protein, of the cell. The principles of the two-point attachment theory and a demonstration of the apparent validity of some of the listed structural requirements for auxin activity are clearly illustrated by a comparison of some of a series of chlorinated phenoxy compounds (Fig. 4.2).

It is necessary to stress that neither the original list of requirements for auxin activity, nor the two-point attachment theory are now accepted as valid. Clearly, the activity of a number of synthetic auxins such as benzoic acid derivatives (Fig. 4.3) cannot be adequately explained on the basis of the list of requirements drawn up in 1938 and the two-point attachment theory, and several alternative hypotheses have been put forward over the years, including "three-point" and "multi-point" attachment theories. The question is still open as to whether the auxin molecule becomes attached to some receptor in the cell by covalent (chemical) bond formation, or by some form of physical association, though the latter is generally considered much more likely. One suggestion, based on studies of the physical properties of active molecules, is that van der Waals and electrostatic forces are important in auxin-receptor association. Thus, a comparison of a range of auxins revealed that molecules active as auxins contain a strong negative charge (arising from the dissociation of the carboxyl group) which is separated from a weaker positive charge on the ring by a distance

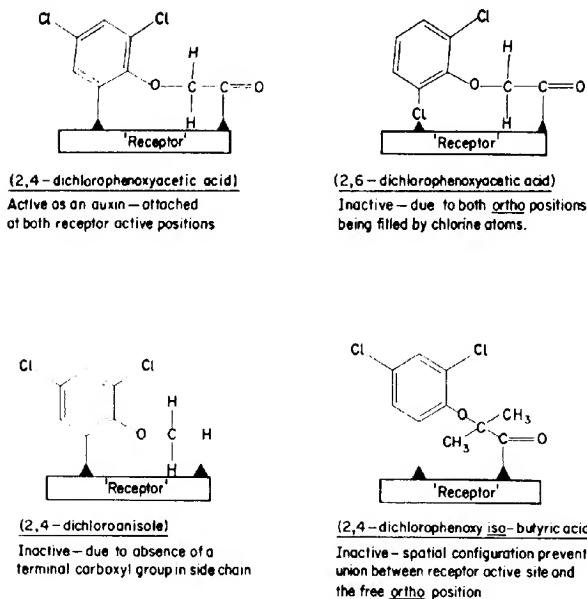


FIG. 4.2. The two-point attachment theory of auxin activity, illustrated by a comparison of 2,4-dichlorophenoxyacetic acid (2,4-D) with three inactive analogues.

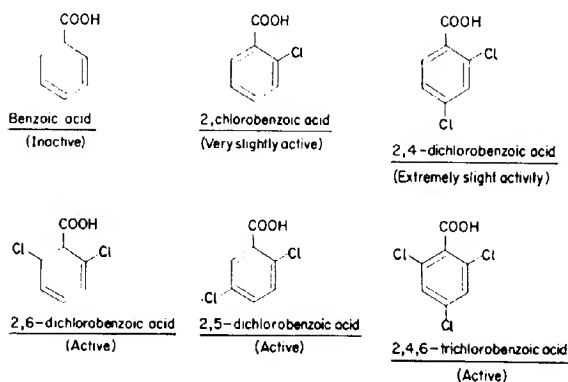


FIG. 4.3. Activity or inactivity as auxins of a series of chlorinated benzoic acid derivatives. Note that 2,6-dichlorobenzoic acid and 2,4,6-trichlorobenzoic acid are active and also have a halogen atom at both *ortho* positions.

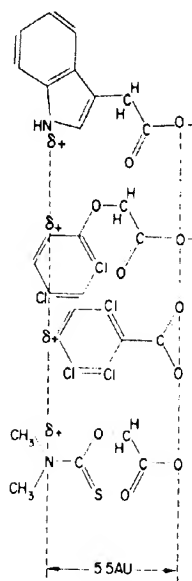


FIG. 4.4. Some diverse molecules active as auxins. They are, however, similar in the possession of a strong negative charge (δ^-) separated from a weaker positive charge (δ^+) by a distance of 5.5 Angstrom units (5.5 AU). From top, indole-3-acetic acid, 2,4-dichlorophenoxyacetic acid, 2,4,6-trichlorobenzoic acid, carboxymethylthiocarbamate. (From K. V. Thimann, *Ann. Rev. Plant Physiol.* **14**, 1-18, 1963.)

of about 5.5 Å (Fig. 4.4), and it has been suggested that this is the essential structural requirement for auxin activity. This hypothesis would explain the relative activities of many synthetic auxins, and the differences in the activity of closely related compounds are apparently due to the effects of substitution in the ring on the position and size of the positive charge. Neither the nature, nor the location within the cell, of the receptor molecule is yet known.

Thus, the intensive study which has been devoted to the molecular requirements for auxin activity has not yet given any clear indication of the basic mechanism by which auxins produce their effects in growth and differentiation. It has, however, led to results of practical importance in the finding of a number of compounds which have proved of enormous value in agriculture and horticulture, such as selective weed-killers, fruit-setting agents and rooting-hormones (p. 135).

Structure-activity relationships of gibberellins. The relationship between molecular structure and biological activity of gibberellins has been less rigorously studied than it has for auxins. Reasons for this are: (i) the shorter time that has elapsed since the discovery of gibberellins

than of auxins, and (ii) it has so far proved impractical to manufacture any "synthetic gibberellins" due to the complex nature of the gibbane carbon skeleton.

As stated earlier, over fifty chemically characterized gibberellins are currently known. All of these have been obtained from natural sources, either from the fungus *Gibberella fujikuroi* or from higher plants. The structures of gibberellins A₁ to A₂₉ inclusive are shown in Fig. 4.5. It can be seen that they are all similar in the possession of the same basic carbon










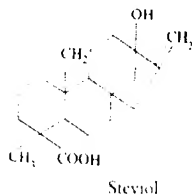
A-Ring	Substituents		C- and D-Rings		
	Carbon position	Group			
	10	-CH ₃	A ₁₂		
	10	-CHO	A ₂₄	A ₁₉	
	10	-COOH	A ₂₅	A ₁₇	
	{ 10 3	{ -CH ₃ -OH }	A ₁₄	A ₁₈	
	{ 10 3	{ -CHO -OH }		A ₂₃	
	{ 10 3	{ -COOH -OH }	A ₁₃	A ₂₈	
	—	—	A ₁₅		
	3; 16	-OH	A ₂₇		
	—	—	A ₉	A ₂₀	A ₁₀
	3	-OH	A ₄	A ₁	A ₂
	2	-OH		A ₂₉	
	3; 1	-OH	A ₁₆		
	3; 2	-OH		A ₈	
	3; 2 12	-OH = O	A ₂₆		
	—	—		A ₂₁	
	3	-OH	A ₇	A ₃	
	4	-CH ₃		A ₅	
	4	-CH ₂ OH		A ₂₂	
	—	—		A ₆	
	—	—	A ₁₁		

FIG. 4.5. A summary showing the range of chemical structures and chemical relationships in the first twenty-nine gibberellins to be isolated and characterized. (Adapted from L. J. Audus, *Plant Growth Substances*, Vol. 1, *Chemistry and Physiology*, 3rd ed., Leonard Hill, London, 1972.)

skeleton. The structural differences between them lie principally in the number and distribution of hydroxyl (—OH) groups, and the degree of saturation of the "A" ring. Present knowledge allows us to assume that for a molecule to behave as a gibberellin it must have a structure similar to that of known naturally occurring gibberellins. On the other hand, a naturally occurring diterpenoid in plants called *steviol*, which does not have the gibbane carbon skeleton, has been found to have some slight growth-promoting properties similar to those of gibberellins. However, this is probably due to conversion of steviol by plant enzymes to an active form, rather than to its having hormonal activity itself.



It should be noted that not all of the known gibberellins are equally effective in stimulating growth. In fact, their activity when tested with different species and varieties of plants can be used as a means of distinguishing between different gibberellins. A good example of this is seen in their effect on the growth of dwarf mutants of maize (*Zea mays*). In maize there are a number of mutant genes, the presence of any one of which results in a dwarf habit of growth. Certain gibberellins have been found to promote the growth of some dwarf mutants, while others are effective with other mutants. It has been suggested that the primary effect of the mutant genes in maize is on the levels of endogenous gibberellins, by interfering with different steps in the biochemical pathway leading to a gibberellin necessary for normal growth. In the case of the d-5-mutant variety of *Zea mays*, it has been found that *iso*-kaurene is formed (instead of kaurene, see Fig. 3.4) and that the plant is unable to metabolize this to gibberellins.

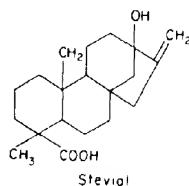
Structure-activity relationships of cytokinins. As we considered earlier in this chapter, the important known natural cytokinins are all substituted adenine compounds that possess side chains of five carbon atoms attached to carbon-6 of the adenine moiety. Systematic studies of many 6-substituted purines have revealed that where the side chain does *not* contain a ring system, the optimum number of side chain carbon atoms is five. Increasing or decreasing the size of the aliphatic side chain reduces physiological activity of the cytokinin, but does not necessarily abolish it completely. Thus, lengthening the side chain to contain as many as ten carbon atoms (e.g. 6-decylaminopurine) does not completely remove activity. A further feature of the side-chain requirements is that the presence of a double bond in the aliphatic side chain usually increases cytokinin activity.

The range of molecules which, though not naturally occurring as cytokinins in plants, do possess cytokinin activity is very large indeed. We have previously mentioned synthetic

cytokinins such as kinetin and benzyladenine (p. 61) that contain ring systems in their side chains. Some of these synthetic cytokinins are at least as, or even more, active than naturally occurring cytokinins. Benzyladenine, for example, is more active than even zeatin (the most potent of the natural cytokinins) in some types of bioassay.

Generally speaking, any modifications made to the adenine ring result in reduction in activity as a cytokinin. Thus, riboside or ribotide derivatives of cytokinins (p. 62) are less active than the free cytokinins, and various other types of attachment or substitution in the adenine ring always reduce, if not actually remove completely, hormonal activity. However, some 1-substituted adenine compounds have been found to have cytokinin activity, but it is possible that this results from enzymatic conversion to the 6-substituted active forms in plant tissues.

Exceptions to the general rule that cytokinins are 6-substituted adenine compounds are seen in a number of phenylureas and the related biurets. Although the phenylureas are very much less active than the subsequently discovered purine cytokinins, several hundred compounds of this type do possess cytokinin activity, among the most active of which is chlorophenylphenylurea. The minimum requirements for activity in the phenylureas are the



—NH—CO—NH— bridge and a planar phenyl ring. At first sight it is difficult to recognize what structural features these compounds have in common with adenine derivatives, but one is the —N—C—N— linkage, of which adenine derivatives have four and the ureas one. The six-membered pyrimidine ring of adenine may be analogous to the phenyl ring of the ureas and active biurets (e.g. fluorophenylbiuret, above) and the amino nitrogen of the purine analogous to a *meta* substituent (e.g. Cl) in phenylureas.

It can be seen, therefore, that studies of the structure-activity relationships of cytokinins have revealed a situation essentially the same as that derived from similar work on auxins, in that a quite bewildering array of compounds are seen to have the capacity to influence growth and differentiation in the manner of cytokinins. This has, so far, prevented any meaningful conclusions being drawn as to a possible receptor site for the initial action of cytokinins.

Structure-activity studies on ethylene analogues. Ethylene, $\text{CH}_2=\text{CH}_2$, is a small unsaturated hydrocarbon, and the physiological activities of a series of ethylene analogues have been

compared with those of ethylene itself. None of the analogues possesses activity as great as that of ethylene, whether the comparison is based on concentrations in parts per million (ppm) or molarity. On a ppm basis, for example, propylene, $\text{CH}_3-\text{CH}=\text{CH}_2$, has activity only about 1/100th that of ethylene; acetylene, $\text{CH}\equiv\text{CH}$, less at 1/2800th (except in the case of the induction of flowering in pineapple, where for unknown reasons acetylene can be as active as ethylene); and allene, $\text{CH}_2=\text{C}=\text{CH}_2$, even less at 1/29000th ethylene activity.

Carbon monoxide, $\text{C}=\text{O}$, has effects on plants similar to those elicited by ethylene, but with only approximately 1/2700th the potency of ethylene itself. Carbon dioxide may act as an antagonist to ethylene action in plants, and this is possibly because CO_2 is a close structural analogue of allene and carbon monoxide but nevertheless lacks certain molecular characteristics which are essential for ethylene action. Thus, CO_2 may compete with ethylene for the active receptor sites for ethylene action in plants, and the physiological response of a tissue to a given concentration of ethylene is determined by, among other factors, the prevailing concentration of CO_2 in the tissue.

The characteristics that a molecule must show to allow it to function, however weakly, as a substitute for ethylene were summarized as follows by Burg and Burg in 1967:

- (1) The molecule must be unsaturated. A double bond confers more activity than a triple bond, and single-bond compounds are inactive.
- (2) Activity decreases with increasing chain length.
- (3) Substitutions that lower the bond order of the unsaturated position by causing electron delocalization reduce biological activity, although steric factors are also important. Thus, the nature of the substituent can affect activity by influencing both electron density in the double bond and overall size and shape of the molecule.
- (4) The unsaturated position must be adjacent to a terminal carbon atom.
- (5) The terminal carbon atom must not be positively charged.

These structural features, derived from studies of the varying degrees of physiological activities shown by ethylene and ethylene analogues, together with the known antagonism to ethylene action shown by carbon dioxide, do not help us to decide the nature of the initial receptor site for ethylene, nor the type of bonding involved in linking hormone and receptor, except that ionic and hydrogen bonding seem unlikely. However, various other experiments have yielded results that indicate that ethylene may be bound to its site of action by means of weak van der Waals forces, rather than by covalent or co-ordination bonding. The effects that ethylene can have on secretion of materials from plant cells lends support to the view that cell membranes may contain the sites of ethylene binding and initial action.

Abscisic acid structure and physiological activity. The natural (+), and synthetic (−), optical enantiomers of ABA have equal activity on plants, but only the 2-*cis* geometric isomer

possesses hormonal activity (see p. 67 for an account of isomerism in ABA). Plant extracts have been found to contain only traces of the 2-*trans* isomer, and it is likely that even these small amounts are formed by isomerization of the natural 2-*cis* isomer during extraction and purification procedures. It therefore appears that only the 2-*cis* form of ABA will fit an unknown receptor site in cells. Studies of ABA analogues have not, so far, been successful in further defining molecular requirements for ABA-like physiological activity, except that the ring double bond appears essential for activity.

Direct Evidence for Hormone Binding to Cell Constituents

The previous section considered how studies of the biological activity of plant hormones and their analogues have not yet yielded any clear idea as to the natures of hormone receptors in plants. Nevertheless, as we saw, activity in each class of plant hormone is determined by structural and often stereospecific properties of the molecules. It is therefore reasonable to consider it probable that there exist receptor molecules able to recognize subtle differences between natural growth hormones and their analogues. Many scientists feel that plant hormone receptors are probably proteins, because of this capacity for recognition of appropriate hormone structure. Such a view is supported by research in animal endocrinology, where several specific proteinaceous receptors for steroid hormones have been isolated and characterized.

None of the attempts made to date have succeeded in unequivocally identifying a plant hormone receptor. Most of such work has concentrated upon the possibility of isolating protein receptors. Various approaches to the problem have been made, mainly using radioisotopically labelled hormones of high specific activity. One method has been to supply the radioactive hormone to plant tissues, and to subsequently homogenize and fractionate the cell constituents by centrifugation or gel filtration, in the hope that radioactivity will be found to be associated with a particular cell fraction. Another has been to apply the labelled hormone to various cell fractions, such as nuclei, isolated chromatin material and membrane fractions, with measurement of the affinity for binding between these and the hormone. Thirdly, attempts have been made to localize radiolabelled hormones within cells by autoradiographic methods. Reasons for the failure to isolate a proteinaceous receptor for any of the plant growth hormones are not at all certain. It should be mentioned that a number of pieces of research have yielded results suggesting that receptor proteins for cytokinins and gibberellins are present in plant cells, but their existence is by no means proven.

It has been recently pointed out by Kende and Gardner that some features of the action of plant hormones indicate that they may be significantly different from animal steroidal hormones with respect to their association with receptor sites, and that it may therefore be incorrect to assume the existence of proteinaceous hormone receptors in plant cells. The dose-response curves for plant hormone action characteristically show that their effects vary over a very wide range of concentration (typically over some four or five orders of

magnitude) which contrasts with the much more restricted effective concentration range for animal hormones. Comparisons of hormone dose-response curves also suggest that whereas the binding of an animal steroid hormone to its receptor fits Michaelis-Menten-type saturation kinetics, the binding of plant hormones do not, which in turn poses the possibility that plant hormone receptors are not specific proteins. If the receptors are not proteins, then one can only guess as to their possible nature. However, experiments referred to below have indicated that the primary point of plant hormone action lies in cell membranes, so that it appears to be a property of certain plant cell membranes to serve as specific receptors for growth hormones. Although one must recognize our current ignorance of the molecular mechanisms involved, it is of course possible that protein receptors may be located in cell membranes.

Mechanism of Action of Auxin in Cell Extension

For the past decade or so two major concepts have existed as to the initial site of auxin action, both of which have been derived from studies of auxin-regulated cell extension growth in coleoptiles, stems and hypocotyls. One of these concepts centres upon the cell wall as the site of auxin action while the other focuses upon effects of auxin on nucleic acid metabolism. To a large extent the work on cell walls and on nucleic acids has been performed by separate research groups, and it is only recently that information and the concepts that have evolved from these parallel studies are beginning to be brought together to formulate a clearer and more detailed picture of the mechanism of action of auxin. A third line of investigation of the problem has been the examination of various effects of auxin on cell membrane properties, and results of these studies, too, are now being related to what is known of the effects of auxin on nucleic acid metabolism and cell wall properties.

Like other developmental processes, cell extension growth involves sequential changes in levels and/or activity of enzymes. In the light of current knowledge of the role of nucleic acids in directing the course of protein synthesis, it seems reasonable to consider the possibility that auxins act by influencing nucleic acid metabolism. As early as 1953 it was reported by Silberger and Skoog that auxin-promotion of growth in tobacco pith callus in aseptic culture was preceded by a proportional increase in RNA level in the cells, and that the maximum measured increases in both RNA and growth occurred in response to the same concentration of IAA. Since this first observation of an effect of auxin on RNA synthesis, numerous other reports have appeared demonstrating positive correlations between growth and auxin-enhanced growth on the one hand, and rates of RNA synthesis on the other. Many of these experiments have involved the use of various inhibitors of protein synthesis. These inhibitors block certain steps in the synthesis of nucleic acids and protein (e.g. actinomycin D which blocks formation of RNA in DNA-dependent RNA synthesis, and cycloheximide which acts at ribosomal level by inhibiting assembly of proteins). For example, Fig. 4.6 shows that the capacity of exogenous auxin to enhance extension growth is inhibited by actinomycin D to the same extent that RNA synthesis is suppressed.

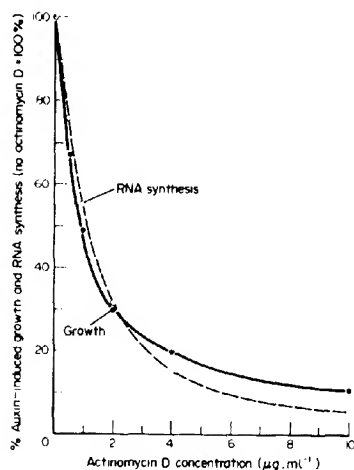


Fig. 4.6. Auxin-induced growth and RNA synthesis are similarly inhibited by various concentrations of actinomycin D. Soybean hypocotyl segments were pretreated for 4 hours with the indicated concentrations of actinomycin D prior to the addition of 5×10^{-5} M 2,4-dichlorophenoxyacetic acid (2,4-D), an auxin. RNA synthesis was measured by incorporation of $(^{14}\text{C})\text{-ADP}$ over the 4-hour growth period. (Adapted from J. L. Key *et al.*, *Ann. N.Y. Acad. Sci.* **144**, Art. I, 49-62, 1967.)

Data such as these certainly imply that continued auxin-produced growth involves RNA and protein synthesis. More direct evidence that auxin can indeed stimulate RNA synthesis has come from experiments using isolated, cell-free, nuclei or chromatin preparations. Such experiments involve incubation of the isolated nuclei or chromatin with RNA-precursors (ATP, CTP, GTP and UTP) that are radioisotopically labelled. Newly synthesized RNA can therefore be measured in terms of incorporated radioactivity. Although available data from these experiments is incomplete, it seems clear that auxin treatment does increase the capacity for RNA synthesis in isolated nuclei and in chromatin preparations. However, these effects of auxin occur only when the cells are exposed to auxin *before* isolation of the nuclei or chromatin. This suggests that the *initial* site of auxin action is located outside the nucleus and that auxin does not act directly at the transcriptional level of control.

It is clear then that auxin can affect enzyme levels through an influence on RNA synthesis (the effect could be to increase total RNA and/or to bring about the formation of qualitatively different RNA). Furthermore, it is possible that auxin could affect enzyme activity more directly and immediately, by affecting enzyme release or activation. By whatever means auxin influences enzyme levels and activities in plant cells, attention is naturally concentrated upon those enzymes that may be thought to be closely involved in cell

enlargement processes. Plant cells are surrounded by a cell wall, and cell growth can occur only if the properties of the wall are changed in such a way as to permit expansion of the protoplast. This basic fact of plant cell growth has logically led to an examination of the possibility that auxin affects wall properties, perhaps by influencing the synthesis of enzymes concerned in cell wall mechanical properties and in wall growth. For the remainder of this section we will consider the relationship between auxin action and the cell wall, and see how studies of this have helped towards a general understanding of the regulation of growth by auxin.

As we saw in Chapter 1, all plant cells, with the exception of those which remain permanently meristematic, pass through two phases in their cycle of growth; these are division and enlargement resulting from vacuolation. The coleoptile of an oat seedling illustrates this pattern in a clear manner, since all cell division ceases when it is about 10 mm in length and all subsequent growth is entirely due to the enlargement of existing cells. Hence, when we study the effects of auxin on the growth of coleoptile sections (p. 48), we are essentially dealing with hormonal effects on cell extension. During cell enlargement, due to vacuolation, irreversible plastic stretching of the cell wall takes place. It is, therefore, tempting to consider that cell vacuolation is a consequence of a softening of the cell wall, for this would inevitably lead to an influx of water into the protoplast for the reasons given in Chapter 1 (p. 5-7). Many experiments have revealed that auxin increases the plasticity of the cell walls. This can be shown by increased plastic deformation of plant organs treated with auxin following the application of a mechanical force (Fig. 4.7). The possible physiological significance of auxin effects on cell wall plasticity is increased by observations that there is a positive correlation between the effects of different auxin concentrations on promotion of elongation growth, and on cell wall plasticity (Fig. 4.8).

During cell enlargement, the cell wall not only stretches, but it also increases in thickness by the deposition of new cell wall material (p. 8). This cell wall growth is stimulated by auxin, and can occur even when cell enlargement is completely suppressed by various means (e.g. by surrounding the tissue with a hypertonic solution of mannitol).

The type of results shown in Fig. 4.7 indicate that cell walls show *viscoelastic* extension, that is, an initial rapid extension followed by a further slower extension (termed "creep"). However, *living cells* may elongate at a *constant* rate for a considerable time and it is thus accepted that growth proceeds by a *series of viscoelastic extensions* driven by the turgor pressure of the cell sap. Moreover, whereas cell walls derived from living tissue treated with auxin give the type of response shown in Fig. 4.8, isolated cell wall material does not respond in this way. This has led to the proposal that auxin does not act directly on the cell wall but rather that it controls certain events in the protoplasm which result in a change in the properties of the cell wall. What are these events which are so influenced? In order to understand this we must consider briefly the structure and physical properties of primary cell walls.

As we saw in Chapter 1 the walls of young growing cells consist of interwoven chains of cellulose microfibrils embedded in a dense matrix of noncellulosic polysaccharides (of several different types) and protein. These components make up approximately 20 per cent by weight of the wall, the remaining 80 per cent being water. In general, one can regard the

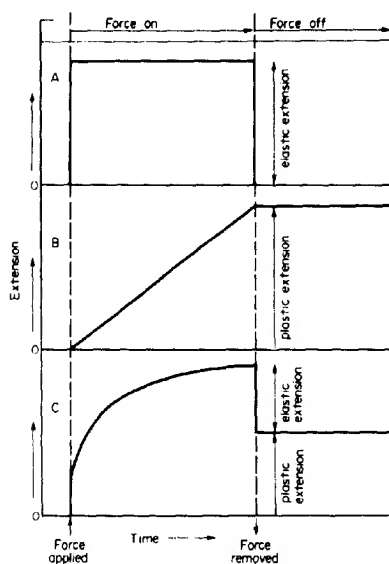


FIG. 4.7. Relationships between extension and time for three contrasting types of materials.

A. An elastic material such as rubber (the molecules of which are bonded together by extensive crosslinks) shows nearly instantaneous extension followed by no further increase with time, but the extension is fully reversible upon removal of the imposed force which caused the extension.

B. Where only few crosslinks or entanglements are present between the molecules, as in short-chain plastics, then irreversible extension occurs by *viscous flow* which is directly proportional to time.

C. In the case of materials such as the primary walls of plant cells, which contain polymers of varying lengths and degrees of crosslinking, the extension is of an intermediate type called a *viscoelastic extension*, in which a certain amount of instantaneous extension occurs initially but is followed by a period of slower and continuous extension at a rate which is nearly proportional to the logarithm of time; the time-dependent component of viscoelastic extension is known as "creep". In fact, the instantaneous part of viscoelastic extension is just the creep which occurs too rapidly to be measured, so that there is no fundamental difference between the two components. Viscoelastic extension may be either partially or completely reversible, depending on the previous history of the material under test. When subjected to mechanical stress for the first time, recovery behaviour is as illustrated in C for plant cell walls—part of the viscoelastic extension can be seen to have been elastic and part plastic in nature. Treatment with auxin increases the plastic component of total extension. Subsequent extensions, provided that the maximum length reached in the first extension is not exceeded, are entirely elastic, and this change in behaviour is termed *mechanical conditioning*. Note, however, that the extension pattern for plant cell walls illustrated in C applies only to *dead* tissues. The walls of *living* cells extend by a continuous series of such viscoelastic extensions, so that extension may proceed at a steady rate for a considerable period of time.

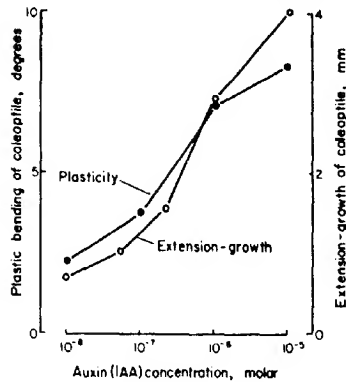


FIG. 4.8. Positive correlation between effects of auxin (IAA) on cell wall plasticity (measured by plastic bending) and on elongation in the oat coleoptile. (From J. Bonner, *Z. Schweiz. Forstw.* 30, 141-59, 1960.)

cell wall as being analogous in structure to steel-reinforced concrete or glass-fibre-reinforced plastic, the cellulose microfibrils acting as the reinforcing element, and the non-cellulosic matrix serving as the stabilizing component. The cellulose molecules within the microfibrils are held together by hydrogen bonding whereas the components of the matrix—both polysaccharides and protein—appear to be connected by covalent bonds. In dicotyledons at least it appears that hydrogen bonding also occurs between microfibrils and matrix and that the matrix polysaccharide involved is a xyloglucan (a polysaccharide with a $\beta 1 \rightarrow 4$ linked glucan backbone as in cellulose but also with frequent xylose side chains and occasional galactose, fucose and arabinose units attached). The cell wall protein is unusual in that in addition to amino acids it contains a very high proportion of the imino-acid hydroxyproline (Fig. 4.9a). Each hydroxyproline unit in the protein is glycosidically connected to an arabinose chain four units long (termed a tetraarabinoside) which is not, however, connected to the rest of the matrix (Fig. 4.9b). The polysaccharides of the matrix appear to be linked to the wall protein via the serine residues of the latter.

As in the man-made structures mentioned above, so in the cell wall the mechanical properties are the resultant of interactions within and between the microfibrillar and matrix components. It follows, therefore, that auxin must in some way affect these interactions. Now it is known that although cell extension requires continued protein and RNA synthesis and respiration, nevertheless if auxin is applied to stem or coleoptile tissue the growth rate increases after a "lag" of a matter of a few minutes (Fig. 4.10) which makes it unlikely that growth is accelerated by changes in the rates of transcription or translation, but rather that auxin is affecting some "pre-formed" system. It is also well established that marked changes occur in the polysaccharides of the wall in response to auxin treatment and indeed, as shown above, it is difficult to conceive how the mechanical properties of the cell wall can

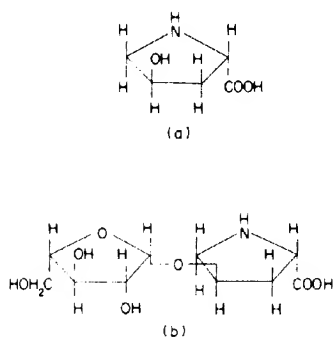


FIG. 4.9.(a) Structure of hydroxyproline. (b) Glycosidic linkage of hydroxyproline to arabinose. Each hydroxyproline unit in cell wall protein is connected in this manner to a tetraarabinose molecule (i.e. four joined arabinose units).

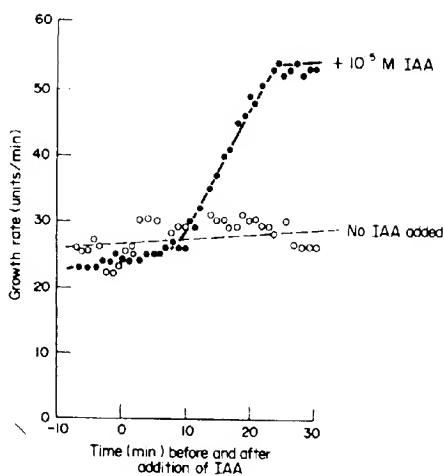


FIG. 4.10. The rate of response of excised pea-stem segments to 10^{-5} M IAA. The elongation growth rate of two batches of segments are shown, and it can be seen that treated segments commenced elongating at a more rapid rate after a lag period of about 10 minutes after application of the auxin. (Adapted from Pauline Penny, *New Zealand J. Bot.* 7, 29-301, 1969.)

be altered without changes of this nature. It seems that since such changes involve the making and/or breaking of covalent bonds then it is likely that enzymes are involved. In the past many workers have attempted to define what changes occur in the polysaccharides of the wall and whether these changes are correlated with variations in enzyme activities within the wall.

These efforts have not on the whole been conspicuously successful, principally because the exact structure of cell wall matrix polysaccharides, and how they were interconnected was unknown. This made interpretation of such work very difficult indeed. However, recent work has given us a much clearer picture of cell wall structure and armed with this knowledge and some new experimental techniques a number of new hypotheses have been advanced. These hypotheses have been greatly influenced by another recent finding, namely that incubation of coleoptile or etiolated stem tissue at low pH (around 3.0) results in extension growth (termed the "acid growth effect") which in the short term at least is similar to that shown by exposure to auxin but without a significant "lag" phase. This led to further work which showed that auxin appears to promote the secretion of H^+ ions (protons) by stem or coleoptile tissue resulting in a lowering of the pH of the wall. The kinetics of the acid-growth effect are very similar to those of growth stimulation shown by auxin-treated tissue. These findings would certainly account for the observed requirement for respiration mentioned above since ion "pumps" and charge separation are features of this process. The hypothetical proton-pump is generally assumed to be located in the plasma membrane (Fig. 4.11).

If we accept that this phenomenon represents the mode of action of auxin in increasing growth rate then three further questions present themselves: (1) How does auxin promote H^+ ion secretion? (2) Why are RNA and protein synthesis required for extension growth? (3) How does changing the pH of the cell wall alter its properties?

The answer to the first question is quite unknown and much further work is needed before a solution is likely to present itself. As regards protein and RNA synthesis the question does not really arise if we distinguish between the *induction* of increased growth and its continuance once induced. Thus, although there is no sound evidence that the induction of growth requires RNA and protein synthesis, continuance of *normal* cell growth requires that the wall must not only change its mechanical properties but much synthesis of new wall material must also occur—this is clearly so because cell walls do not become thinner as they extend. Thus a continuous supply of biosynthetic and hydrolytic enzymes is necessary and this probably involves both RNA and protein synthesis. Indeed it appears that the amount and/or activity of such enzymes—cellulose synthetase for example—is increased by auxin treatment, although this occurs some time after the promotion of growth; conversely, such changes are often suppressed if growth is inhibited even though auxin is present.

No firm answer is yet available to the third question but clearly H^+ ions could act in two obvious ways, namely by breaking acid-labile bonds directly or by making conditions more favourable for various enzyme mediated modifications of the wall, e.g. by changing the pH of the wall so that it is nearer the pH optimum of some critical enzyme(s). Thus, it

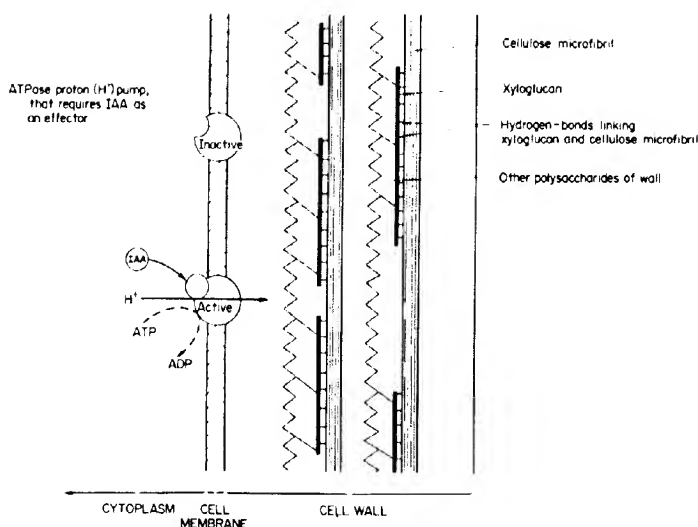


FIG. 4.11. Diagram to illustrate the hypothetical scheme of auxin (IAA) action as an effector of a membrane-bound proton (H^+) pump. The pump is envisaged as containing ATPase, and being active in pumping H^+ ions from the cytoplasm into the cell wall only when IAA is present. The secreted H^+ ions could cause breakage of hydrogen bonds that link together xyloglucan polymers and cellulose microfibrils. When the wall is under tension (cell turgor pressure) this would result in a creep of the xyloglucan along the microfibril and thus give wall extension and cell enlargement. (Adapted from P. J. Davies, *The Botanical Review*, 39, 139-71, 1973.)

has been suggested that the bond linking the non-cellulose polysaccharides and the cell wall protein may be directly broken at low pH, but no firm evidence has been adduced to this end. If the process is enzyme controlled, the xyloglucan component of the matrix in dicotyledons may be the critical component. It has been shown auxin influences the "turn-over" of xyloglucan more markedly than it does of other polysaccharides. That is, both breakdown and synthesis of this molecule are accelerated by auxin. Moreover, this effect occurs very rapidly in response to auxin treatment. As we showed above, the xyloglucan appears to interact with the microfibrillar component of the wall and a change in this interaction would be expected to affect wall properties. At present, much work is being done to identify the enzyme(s) responsible for this increased turnover.

Finally it should be emphasized that whereas the above hypothesis is feasible for dicotyledons, monocotyledons do not appear to possess xyloglucans and hence another mechanism is called for in their case.

Thus, the mechanisms involved in plant cell wall growth are gradually being elucidated, and the role of auxin in the overall process is beginning to be understood. However, it should be remembered that although cell enlargement is in many respects the most charac-

teristic result of auxin action, it is by no means the only one, nor even necessarily the first to appear. Thus, auxin can induce rapid increases in the rates of respiration and of protoplasmic streaming. Also, a number of responses to auxin do not immediately involve cell vacuolation (e.g. cambial division, root initiation, and correlative inhibition of axillary buds). In other words, although auxin may induce a rapid increase in the rate of cell-wall loosening, this does not necessarily represent the primary, or only, point of auxin action. Finally, it should be mentioned that not only auxins, but also gibberellins induce cell-wall loosening in situations where gibberellins promote cell elongation. It is only recently that this wall-loosening effect of gibberellin has been unequivocally demonstrated, and very little work has yet been done to compare auxin and gibberellin effects on cell-wall growth.

Mechanism of Action of Gibberellins

Research into the mode of action of gibberellins has been greatly helped by discoveries that the levels of activity of certain specific enzymes can be affected by the amount of gibberellin present. Enzymes whose activities are increased by gibberellins include α - and β -amylase, protease, and ribonuclease in germinating barley seeds; and nitrate reductase and ribulose diphosphate carboxylase in leaves of some species. In sugar-cane stems, on the other hand, gibberellins appear to inhibit synthesis of invertase and peroxidase. These regulatory effects of gibberellins on known enzymes have permitted studies of their hormonal action in very much simpler systems than that of cell extension growth which has served as the model for auxin action. Of particular interest is the question whether gibberellin regulation of the level of enzyme activity results from a specific alteration of RNA-directed protein synthesis.

The most studied example of an enzyme whose level of activity can be controlled by gibberellin is α -amylase in barley seed. As we consider later (p. 278), α -amylase is not present in the dry, unimbibed, barley seed, but appears in and is secreted from aleurone layer cells in response to gibberellin transmitted from the germinating embryo. Aleurone tissue isolated from ungerminated barley seed contains only traces of α -amylase activity, but incubation of aleurone pieces in gibberellin solutions results in a very marked increase in the amount of α -amylase activity after a lag phase of at least 8 hours (Fig. 4.12). Stimulation by gibberellin of the appearance of α -amylase activity is prevented if inhibitors of the synthesis of RNA and protein are included in the incubation medium of the aleurone layers, which suggests that gibberellins may regulate α -amylase activity through effects on RNA synthesis. Also, an inhibitor of RNA synthesis such as actinomycin-D has greatest inhibitory effect during the first few hours after gibberellin is added, whereas a protein synthesis inhibitor such as cycloheximide continues to inhibit the appearance of α -amylase activity after the initial lag phase. Results such as these indicate that for α -amylase to appear in the aleurone cells in response to gibberellin, conditions must permit the process of RNA synthesis during the lag phase, but that thereafter α -amylase synthesis can continue using the RNA already formed as template (i.e. the α -amylase m-RNA appears to be long-lived).

Evidence that the α -amylase induced by gibberellin arises as a result of *de novo* enzyme

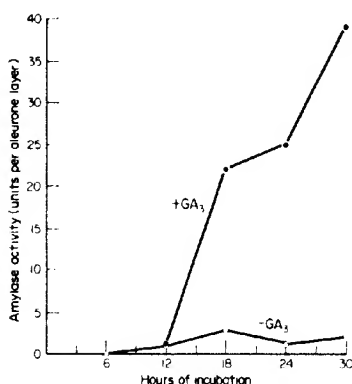


FIG. 4.12. Time course of enzyme (α -amylase) release from isolated barley aleurone layers incubated in media containing 10^{-6} M GA₃ (+ GA₃) or no gibberellin (- GA₃). Not until layers after 8 to 12 hours incubation do measurable quantities of enzyme appear, and the stimulatory effect of gibberellin become clear. (From K. M. Bailey, I. D. J. Phillips and D. Pitt, *J. Exp. Bot.* 27, 324-36, 1976.)

synthesis was provided during the late 1960s. In 1967 Filner and Varner provided conclusive evidence that *all* the α -amylase produced in response to GA₃ treatment is synthesized *de novo* from amino acids. They obtained this proof of the effect of GA₃ on enzyme synthesis by use of *density-labelling* techniques. Barley aleurone layers were incubated with GA₃, together with either normal water (H₂O¹⁶) or water containing the heavy isotope of oxygen (H₂O¹⁸). The natural storage proteins in the aleurone cells were thus hydrolysed during protease action in the presence of either H₂O¹⁶ or H₂O¹⁸, resulting in the formation of O¹⁶-containing or O¹⁸-labelled amino acids. The latter, heavy-isotope containing, amino acids are said to be density labelled, and proteins formed from them will also be density labelled (i.e. they will be heavier, or more dense) than proteins formed from O¹⁶-containing amino acids. Following ultracentrifugation of enzyme extracts, Filner and Varner found that α -amylase formed in GA₃-treated aleurone cells incubated with H₂O¹⁸ was of about the theoretically expected 1 per cent greater density than α -amylase from H₂O¹⁶-incubated aleurone cells (Fig. 4.13), which demonstrated that all of the induced α -amylase was newly synthesized from amino acids during incubation with H₂O¹⁸.

There seems little doubt, therefore, that gibberellins are able to stimulate the synthesis of a new RNA species (perhaps a messenger-RNA) that is required for the formation of an enzyme such as α -amylase. However, it is now known that GA₃ stimulates the level of α -amylase activity in barley aleurone cells *before* a rise occurs in the rate of RNA synthesis, which suggests that the *first* responses of aleurone cells involve the release of preformed enzyme and that only after this has occurred does the stimulatory effect of GA₃ on the synthesis of α -amylase become important. In fact, it is now generally considered that the earliest effect of gibberellins in the barley aleurone system is to influence various membrane

systems already existing in the cells. The effects of gibberellin on membranes of the aleurone cells have been found to include: (a) an increase in membrane synthesis (particularly rough endoplasmic reticulum), (b) stimulation of formation of hydrolytic enzyme-containing vesicles and microbodies from the endoplasmic reticulum, and (c) promotion of the secretion of α -amylase out through the plasma membrane. In addition, experiments on "model" (artificially made) membrane systems have revealed that GA_3 increases their permeability to both uncharged molecules such as sugars and to inorganic ions.

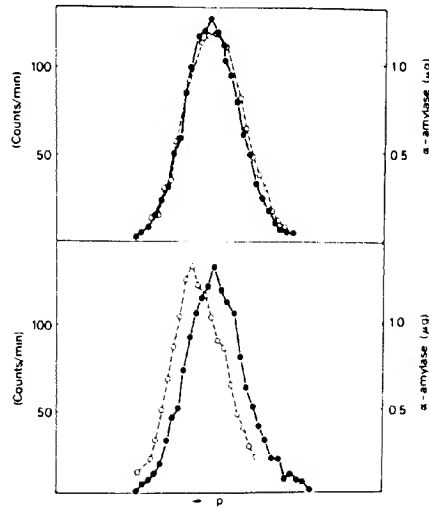


FIG. 4.13. Evidence from "density-labelling" that the entire α -amylase molecule is synthesized *de novo* from amino acids in barley aleurone cells in response to gibberellin treatment. The graphs show the distribution of α -amylase on a CsCl density gradient (ρ = density). Above: Coincidence of densities of α -amylase formed in presence of H_2O^{18} (○---○) and tritiated (3H) marker α -amylase (●—●). Below: Greater density of α -amylase formed in presence of H_2O^{18} (○---○) compared with the marker α -amylase of normal density (●—●). (Adapted from P. Filner and J. Varner, *Proc. Nat. Acad. Sci.*, **58**, 1520-6, 1967.)

Taken all together, these various observations indicate that early effects of gibberellins involve quantitative and qualitative changes in certain membranes and membrane systems in the cell. It seems likely that at least some of these effects on membranes precede the stimulation of RNA and protein synthesis. For example, synthesis of the important membrane phospholipid, lecithin, is increased with 2 hours of GA_3 treatment in aleurone cells. Thus, the mechanism of action of gibberellins appears to be similar to that of auxins insofar as both these categories of growth hormone appear to act first as activators of some preformed, perhaps membrane associated, system, following which longer-term regulatory effects occur through alteration of RNA and protein synthesis.

Mechanism of Action of Cytokinins

Cytokinins appear to play important roles in the regulation of cell division and all known endogenous cytokinins are purine derivatives (primarily of adenine). Thus, research into their mechanism of action has tended to concentrate upon their relationships with nucleic acids. No clear picture has yet emerged, however, although, like auxins and gibberellins, cytokinins quite clearly do have the capacity to stimulate RNA and protein synthesis in plant cells. Some workers have reported that all fractions of RNA (m-RNA, r-RNA and t-RNA) are increased after cytokinin treatment, but others have found that only r-RNA levels are raised.

From the mid-1960s much interest was centred upon the possibility that cytokinins may exert their hormonal effects through modification of specific transfer-RNAs, following the discovery that cytokinin groups occur in certain species of t-RNA. In both serine t-RNA and tyrosine t-RNA adenine nucleotides occur which possess side chains which are isomers of those of the most hormonally active cytokinins. Furthermore, in each case the substituted adenine moiety of the cytokinin was found to be located immediately adjacent to the anticodon of the t-RNA (Fig. 4.14). It was realized that a 6-substituted purine (all natural cytokinins are of this chemical nature) positioned adjacent to the anticodon loop would permit maintenance of the correct spatial arrangement of the t-RNA, and so preclude the possibility of an incorrect triplet of nucleotides being recognized by the codon of m-RNA

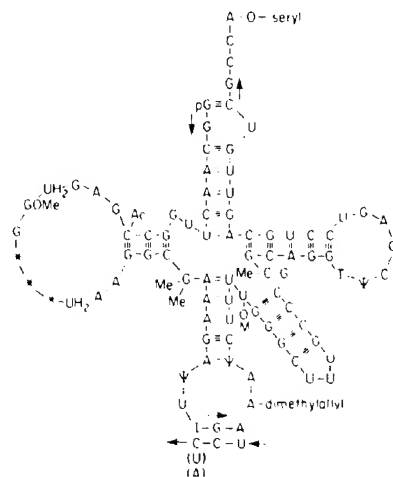


FIG. 4.14. Structure of t-RNA for serine (isolated from yeast cells), showing the location of the cytokinin dimethylallyladenine adjacent to the anticodon. A = adenine; C = cytosine; G = guanine; T = thymine; U = uracil. (From H. G. Zachau *et al.*, *Angew. Chem.* **78**, 392, 1966.)

on the ribosome. In other words, it seems likely that the presence of t-RNA in a molecule of the cytokinin type is essential for normal codon-anticodon interaction between m-RNA and t-RNA on the ribosome. The hypothesis that cytokinins exert their regulatory effects through t-RNA functioning at translation therefore became very attractive.

However, this hypothesis has more recently received severe criticism on a number of grounds. For example, in the normal biosynthesis of t-RNA modification of component bases probably occurs after the primary structure of the polynucleotide has been established, which means that the characteristic side chain on carbon-6 of the adenine moiety of an incorporated cytokinin would be attached *after* the adenine portion is incorporated. This would make it impossible for substances such as kinetin, zeatin, etc., to be incorporated intact into t-RNA. Other evidence against cytokinins acting through their being incorporated into t-RNA includes the finding that t-RNA of *Zea mays* seed contains *cis*-zeatin, whereas the naturally occurring cytokinin in the same seed is *trans*-zeatin, which makes it difficult to believe that the cytokinin is a precursor in t-RNA synthesis. Particularly convincing evidence against the t-RNA hypothesis for cytokinin action has come from studies of rates of incorporation of radioisotopically labelled cytokinins into t-RNAs of cells responding to the hormone. Although the literature on the subject is rather conflicting, in general it has been found that the quantitative limits of cytokinin incorporation into t-RNA are far too low to regard the process as being the basis for the regulatory effects of cytokinins. Moreover, [^{14}C]-6-benzylamino-9-methyl-purine, although active as a cytokinin, does not become incorporated into t-RNA at all due to the masking of the carbon-9 position by a methyl group.

Other, more recent, lines of research are tending to suggest that cytokinins may act as direct regulators of enzyme activity, rather than of enzyme synthesis. Enzymes whose activities are influenced by cytokinins include respiratory kinases, particularly pyruvate kinase. In view of the well-known regulatory effects of cyclic-AMP on kinases in mammalian cells, and some reports that addition of cyclic-AMP to plant cells can induce responses similar to those elicited by cytokinins, it has been suggested that either, (a) cytokinins act in some way equivalent to that of cyclic-AMP in mammalian cells or (b) that cytokinin effects are mediated through a cyclic-AMP system in plants. However, it must be emphasized that very little factual evidence exists to support any of these concepts which therefore remain highly speculative.

The Mechanism of Action of Ethylene

As with the other categories of plant growth hormones, a number of possible initial sites of action for ethylene have been suggested, none of which appear to explain adequately the diverse effects of this substance on the physiology of plants.

Because exposure of plant tissues to ethylene can result in quantitative and qualitative changes in enzymes, one possibility is that ethylene may regulate RNA-directed protein synthesis. Various enzymes have been found to increase in activity following the addition

of ethylene, e.g. cellulase, peroxidase, phenylalanine ammonia lyase, and phosphatase. The stimulatory effect of ethylene on cellulase activity is particularly pronounced in the separation layer cells during leaf abscission (Chapter 12). At least part of this increase has been established as resulting from ethylene-induced *de novo* synthesis of cellulase. However, promotion of leaf abscission processes by ethylene has been found to occur too rapidly to be wholly explained by a mechanism necessarily involving enzyme synthesis, and it has, in fact, been shown that a major effect of ethylene is to cause the rapid release of cellulase from a bound form and its secretion through the plasma membrane into the cell walls of the separation layer cells. Similarly, ethylene enhances the secretion of α -amylase from barley aleurone layer cells.

These rapid effects of ethylene on secretory processes, taken together with observations that some ethylene responses cannot be inhibited by actinomycin D or cycloheximide, and that growth rates can be altered within 5 minutes of exposure of a tissue to ethylene, make it clear that ethylene is able to regulate developmental activities by mechanisms that do not immediately depend on protein synthesis. Thus, it is possible that ethylene affects the activity of the "proton pump" which appears to be involved in cell extension (p. 87).

In view of the high lipid solubility of an olefine such as ethylene, one at least superficially attractive hypothesis of ethylene action proposes that the receptor site for ethylene may be located on the surface of one or more types of cell membrane, and that an association of ethylene with membranes would influence their permeability properties. Although evidence is indeed accumulating that ethylene can regulate secretory phenomena and the transport of materials through plant cell membranes, these effects appear to be mediated by a more sophisticated mechanism than simple physical effects of ethylene on the permeability of membranes.

It is only realistic, therefore, to recognize that the mechanism of action of ethylene probably includes both short-term rapid effects on cell membranes and longer-term effects on nucleic acid and protein metabolism, but that we have very little idea of how these come about.

The Mechanism of Action of Absciscic Acid

Absciscic acid (ABA) resembles the other plant growth hormones in that when applied to plant cells it may elicit changes in the levels of activity of various enzymes, and in the pattern of nucleic acid metabolism. Thus, for example, ABA suppresses the appearance of α -amylase activity in barley aleurone cells.

In the same way that the mechanism of action of the other growth hormones has been sought in terms of a direct modification of nucleic acid metabolism and protein synthesis, so too have the effects of ABA on these processes been investigated. Several possible ways have been suggested in which ABA could suppress levels of enzyme activities. Findings that ABA enhances ribonuclease (RNase) activity led to suggestions that it is this effect of ABA which leads to lower RNA levels and a consequent fall in the rate of protein synthesis.

However, some subsequent work indicated that ABA can reduce total RNA synthesis within 3 hours of its application, but that RNase activity does not increase until after 8 hours (Fig. 4.15), which suggests that the first effect of ABA is not on RNase synthesis or activity.

The means by which total RNA synthesis is reduced in some tissues by ABA has not yet been elucidated. Some experiments have indicated that chromatin activity (i.e. available template) is reduced by ABA, and others that reduced RNA synthesis in the presence of ABA is attributable to a fall in the level of activity of RNA polymerase.

In contrast to the situations where ABA clearly causes a reduction in total RNA content of cells, it has been found that in barley aleurone tissue exogenous ABA is able to inhibit α -amylase synthesis but without affecting *total* RNA or protein synthesis. Furthermore, the inhibitory effect of ABA on α -amylase synthesis in the aleurone cells does not appear to occur through an inhibition of synthesis of the m-RNA for α -amylase. For these and other reasons, it has been proposed that ABA influences either the synthesis or the activity of a regulator RNA that is required for the translation of the α -amylase m-RNA in barley aleurone tissue.

Despite the undoubted influence of ABA on the synthesis of proteins, however it takes place, a number of the known effects of ABA are now known to occur too rapidly to be explained in these terms. Closure of stomata in response to ABA has been measured within a minute of time of application of the hormone, and the prevention by ABA of activation

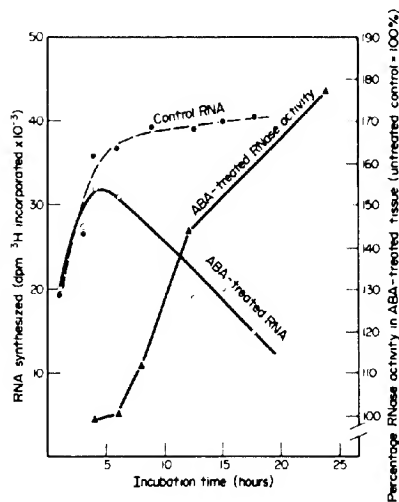


FIG. 4.15. Time-course of effects of 3.8×10^{-6} M ABA on total RNA synthesis (expressed as d.p.m. ^3H -labelled cytidine incorporated in RNA per gramme fresh weight of tissue) and on total RNase activity, in *Zea mays* coleoptile cells. (Adapted from J. H. M. Bex, *Planta*, **103**, 1-10, 1972.)

of the enzyme phosphorylcholine glyceride transferase also occurs too rapidly to be explained by a mechanism that involves effects on RNA synthesis. Furthermore, auxin-dependent elongation growth in coleoptiles can be inhibited within a few minutes of time of addition of ABA. Such observations of rapid effects of ABA are similar to what has been observed for the other plant growth hormones, and suggest that at least some of the regulatory effects of ABA are independent of nucleic acid-directed protein synthesis.

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CHAPTER 5

Hormonal Control in the Whole Plant

IN THE preceding chapter we discussed the possible modes of action of the major groups of growth substances at the molecular and cellular levels and we now have to consider their role in the control of growth and differentiation at the tissue, organ and whole-plant levels. The orderly series of changes so characteristic of development clearly require control systems to ensure that there is co-ordination of growth and differentiation in both *space* and *time*. Thus, the development of a leaf primordium is accompanied by the differentiation of vascular tissue in the neighbouring stem tissue (p. 34). Similarly, the growth of an embryo following fertilization is normally accompanied by growth of the surrounding tissues of the ovule. These are examples of the phenomenon of *correlation*, which we shall consider in more detail later.

Correlation of growth in different regions is also seen at the whole-plant level. The indeterminate pattern of growth of higher plants by apical meristems leads to the steady accretion of mature tissue in the older parts of the plant body, while meristematic activity is maintained at the shoot and root apices. However, potential meristematic regions still remain in the older parts, such as the cambium, axillary buds, developing seeds and fruits, and storage organs. Hence there is a need for control systems to co-ordinate growth in the various regions of the plant, which we shall refer to as *spatial co-ordination*.

There is also a need for *temporal co-ordination* of development, which involves an orderly sequence of changes at all levels of organization. Thus, the whole plant passes through a succession of phases during its life cycle, viz. germination and vegetative growth, flowering and fruiting, ripening and senescence and dormancy. This orderly sequence of changes must also involve control systems which ensure co-ordination in *time*.

A control system may operate spontaneously within the plant itself, i.e. its action may be *autonomic* in origin, or it may be activated or modulated by *environmental* factors. Since environmental conditions do not normally vary appreciably around the different parts of the shoot system, spatial co-ordination within the plant is, in general, achieved by autonomic control systems, whereas environmental factors are frequently important in temporal co-ordination.

Autonomic control is expressed at both the *intracellular* and the *intercellular* levels of

organization. Control of development at the intracellular level is reflected in the appearance and disappearance of activities of different enzymes during various phases of cell growth and differentiation, and this is achieved in several ways, including regulation of nucleic acid and enzyme synthesis and the activation and inactivation of preformed enzymes. Thus, intracellular control of developmental processes quite clearly involves regulation of gene activity. But intercellular control mechanisms must also, of course, ultimately be determined by the genotype.

Spatial co-ordination in plants appears to depend upon the movement of substances between cells and tissues. Such movement may be either (1) *short range*, between adjacent or neighbouring cells or (2) *long range*, involving relatively long-distances interactions. Short-range interactions, involving the movement of protein molecules, appear to be involved in pollen/stigma recognition reactions (p. 326), but whether they also are important in interactions between somatic plant cells is problematical. On the other hand, there is considerable evidence that plant growth-regulating substances play a vital role in intercellular interactions.

Growth substances have been demonstrated to perform various important functions in growth and differentiation, particularly those (a) in which relatively long-distance correlative control is exerted by one organ or region on another, and (b) where environmental effects are apparently mediated through modulation of internal growth substance levels and distribution within the plant body. There is much more evidence for the role of hormones in the control of growth and differentiation in *existing* organs, than for their possible role in the initiation of tissues and organs. Nevertheless, there is a possibility that the major groups of growth substances do play a role in determining the sites of initiation of tissues and organs. Thus, it may be significant that the known growth substances can induce the formation of roots on shoot cuttings and the initiation of buds and roots in callus cultures (Chapter 6).

In order to fall within the traditional definition of a hormone, a substance must be released from the cells in which it is formed and produce an effect in other cells, i.e. the sites of production and action must be separate and movement of the hormone is required. Moreover, to effect control of a process the hormone must be capable of modulation in space or in time. These criteria are certainly met in the hormonal control of growth of the *Avena* coleoptile, where auxin produced in the tip stimulates cell extension in the base, and we shall meet other examples of control by auxin. However, it is more difficult to demonstrate that the other main groups of plant growth substances conform strictly to the traditional definition of a "hormone".

It is clear that the transport and distribution of growth substances within the plant must play a crucial role in their function in spatial co-ordination, and much attention has been devoted to elucidating (a) the sites of biosynthesis, (b) the patterns of hormone movement and distribution from these sites, and (c) the manner in which various environmental factors, such as light and gravity, affect hormone levels and distribution in the plant. Hence we shall first consider some of these latter topics, and then discuss the evidence for hormone control and co-ordination in various aspects of development. In later chapters we shall see

that growth substances probably also play an important role in temporal co-ordination in growth responses to environmental factors, such as daylength and temperature.

THE TRANSPORT OF PLANT GROWTH HORMONES

From their sites of synthesis, growth hormones are transported to other regions of the plant, influencing the cells and tissues with which they come into contact. One would reasonably expect, therefore, that the translocation of these substances would be strictly regulated. Nevertheless, as we shall consider below, although a limited amount of evidence exists to suggest that the transport of abscisic acid, cytokinins and gibberellins may normally follow particular patterns, only auxin translocation has been unequivocally demonstrated to be polarized (i.e. auxins are usually transported along the longitudinal axis of the plant more rapidly in one direction than in the opposite direction). The polar nature of auxin transport is undoubtedly of great importance in the co-ordination of growth and differentiation in different regions of the whole plant. For this reason we will deal first with what is known of auxin transport in plants, following which the transport of the other growth hormones will be considered.

In the shoot tissues which have been studied (coleoptiles, stems, hypocotyls, petioles and flower-stalks), auxin moves more rapidly *basipetally* (i.e. from morphologically apical to more basal regions) than *acropetally* (from basal to apical regions). As we shall consider later, auxin transport in roots also appears to be polar, but there is evidence that the preferred direction of transport may be either acropetal or basipetal, depending on the region of the root.

Auxin Transport in Shoot Tissues

Polar basipetal auxin transport occurs in all organs of the vegetative shoot. The majority of experiments which have shown this have been conducted with short excised segments (usually 5–10 mm long) of coleoptiles, stems, petioles, etc. In principle, the technique is to apply an auxin to one end and to follow its movement along the segment. Various methods have been adopted to determine how much auxin has been transported, and how far, in such segments, but most commonly a “donor–receiver” agar block system has been employed. In this, an agar block containing auxin (the “donor block”) is placed against one cut end of a segment of tissue, and another agar block (the “receiver block”) against the opposite end. Auxin molecules enter the segment from the donor block, are transported through the segment and eventually emerge into the receiver block. Once auxin starts to enter the receiver block, its concentration there rises linearly with time under carefully controlled experimental conditions. The intercept on the time axis of the straight line of increase in auxin content of the receiver block provides an estimate of the average time taken for

auxin molecules to pass from one end of the segment to the other (Fig. 5.1A). Since the length of the segment is known, auxin movement can be expressed in terms of velocity (distance moved in unit time).

Using the donor-receiver block method, Went, in 1928, found that auxin moved only basipetally in *Avena* coleoptile segments. Irrespective of the orientation of a segment with respect to gravity, auxin appeared only in a receiver block placed against the morphological basal end with a donor block positioned at the morphological apical end. The auxin used in Went's experiments was unknown, but it was collected from *Avena* coleoptile tips and was probably IAA. Other investigators repeated Went's experiment, and confirmed the existence of polar basipetal IAA transport in coleoptiles, stems, hypocotyls and petioles. Earlier workers, like Went, measured the quantity of IAA in receiver blocks by bioassay. More recently, the availability of radioactive auxins has allowed more precise experimentation, and this has revealed that (a) auxin transport in aerial organs is not exclusively polar, for some acropetal as well as basipetal movement takes place (Fig. 5.1B), and (b) in addition to IAA, certain synthetic auxins, such as 2,4-dichlorophenoxyacetic acid (2,4-D), indole-3-butyric acid and naphthalene acetic acid (NAA) are transported in a polar manner.

The velocity of basipetal polar auxin transport has been measured in various organs by a number of workers. Values obtained for IAA polar transport all lie between 5 and 15 mm per hour. Synthetic auxins, although transported in a polar manner, apparently move more slowly. For example, 2,4-D moved basipetally in *Phaseolus vulgaris* petiole segments at a velocity of only 1 mm per hour, whereas the equivalent figure for IAA was 6 mm per hour.

The velocity of acropetal auxin transport has not been rigorously determined for aerial organs, but it is normally very much lower than that of basipetal transport. However, the differential between the velocities of basipetal and acropetal auxin transport is influenced by a number of factors. Thus, polarity of auxin movement declines with increasing age of the transporting tissue. It is not yet clear whether this is due to a decrease in basipetal transport, or an increase in acropetal transport, or both. There is, nevertheless, no doubt that maturation processes in a tissue are associated with a gradual reduction in the polarity of auxin transport. Because of this, it has been suggested that polar transport of auxin occurs only in association with cell elongation. However, careful experiments by McCready and Jacobs in 1967 showed that, in bean petiole segments, basipetal polar transport of 2,4-D was less when the segments were elongating rapidly in the presence of gibberellic acid (GA_3) than when their growth was inhibited by mannitol. On the other hand, earlier experiments demonstrated that GA_3 stimulated basipetal IAA transport in stem tissues. Thus, we know neither the significance nor the basis of reduced polarity of auxin transport in mature tissues.

Gravity appears to have some influence on basipetal polar transport of auxins, for several workers have found that when a normally erect organ is placed horizontally, or inverted, then the velocity of basipetal auxin transport is reduced. This phenomenon may be involved in geotropic responses of plant organs, although much more work needs to be done to evaluate this possibility.

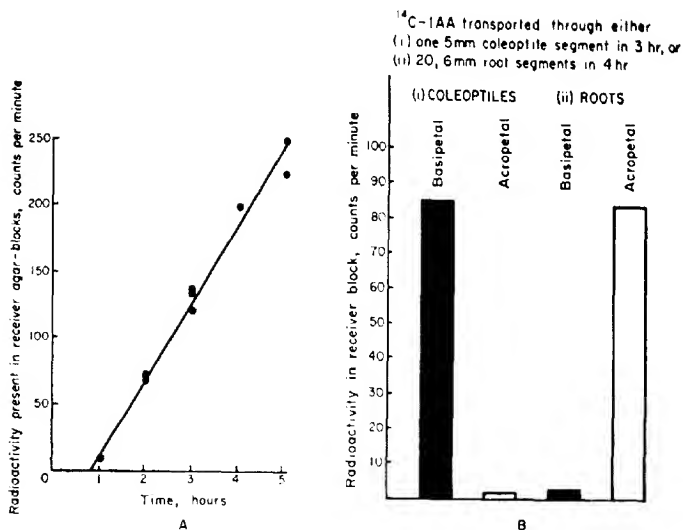


FIG. 5.1. A. Estimation of the velocity of basipetal polar transport of indole-3-acetic acid (IAA) in bean petiole segments. Radioactive IAA (^{14}C -IAA) at a concentration of $50 \mu\text{M}$ in agar-gel (the "donor block") was supplied to the apical end of each segment, and a blank agar "receiver block" was placed against the basal end. Radioactivity appearing in the receiver block was determined at hourly intervals. The graph line showing radioactivity present in the receiver block intercepts the time axis at 0.8 hour. The petiole segments were 5.44 mm long, which means that IAA was transported basipetally at velocity of 6.8 mm/hour. (From C. C. McCready and W. P. Jacobs, *New Phytol.* **62**, 19-34, 1963.)

B. Polar transport of indole-3-acetic acid (IAA) in coleoptile segments (i), and in root segments (ii) of *Zea mays* c.v. "Giant White Horsetooth". Donor agar-blocks containing radioactive IAA (^{14}C -IAA) were placed on either the apical end (basipetal transport) or basal end (acropetal transport) of coleoptile or root segments. The amount of radioactivity (counts per minute) which appeared in blank agar receiver blocks at the opposite ends of the segments was determined. In coleoptiles polar auxin transport is basipetal, but in roots the direction of movement is acropetal. (Coleoptile data from M. B. Wilkins and P. Whyte, *Planta (Berl.)* **82**, 307-16, 1968; root data from M. B. Wilkins and T. K. Scott, *Nature*, **219**, 1388-9, 1968.)

Despite detailed studies of polar auxin transport, extending over 40 years, we still do not know the pathway of auxin movement. The velocity of polar auxin transport (0.5-1.5 cm per hour) is very much less than that of solute movement in the phloem (10-100 cm per hour), and the direction of solute flow in the phloem in the upper stem is acropetal rather than basipetal. For this and other reasons, it does not seem likely that auxin normally travels in the phloem. The other vascular tissue, the xylem, clearly does not usually serve as a transporting channel for auxin, for here again the flow is upwards, and dead xylem elements

would be unable to provide the energy required for polar auxin transport. Early work indicated that *all* cells in coleoptile segments are capable of transporting auxin basipetally at 1 cm per hour. We cannot be certain that this is true for stems, for *Coleus* stem pith-segments failed to transport IAA at all unless a strand of vascular tissue was present. In fact, a number of workers have found over the past few years that basipetal auxin transport in stems occurs primarily or solely in tissues of the vascular strands. Although by no means certain, it does appear likely that, in stems at least, the procambium, cambium and newly formed derivatives of the cambium (particularly phloem initials) may provide the principal routes for polar auxin transport. In coleoptiles, on the other hand, several workers have reported that basipetal polar auxin transport takes place through the non-vascular parenchyma at least as readily as through the vascular tissues.

Auxin Transport in Roots

Until relatively recently, very few direct studies were made of auxin movement in roots, and perhaps because of this, a great deal of confusion existed over this matter for many years. However, experiments by several groups of workers since 1964 with root segments, using radioactive IAA and the donor-receiver agar block method, have amply confirmed that roots of a range of species show polar transport of auxin, and that the direction of movement along most or all of the root is *acropetal* (Fig. 5.1B). This is the reverse of the situation in shoot tissues, and the full physiological significance of the difference with respect to the normal regulation of root growth and geotropism remains to be evaluated. The velocity of polar acropetal auxin transport in roots has been found to be approximately 1 cm per hour, which is similar to that of basipetal polar auxin transport in shoot tissues. Despite the evidence for acropetal movement along most of the root, some recent studies have indicated that auxin transport may show basipetal polarity in the more apical parts of roots, but again the possible physiological significance of such a situation is obscure at present.

As is the case for stem tissues, available experimental evidence suggests that auxin transport in roots occurs primarily in tissues of the vascular strands, particularly the cambium and newly formed phloem.

The Mechanism of Polar Transport of Auxin

Polar transport of auxin in plant organs is a manifestation of the existence of polarity in each individual cell (Chapter 13). Experiments with segments of organs such as coleoptiles have shown that the polarity of transport (i.e. the ratio, basipetal transport/acropetal transport) increases approximately exponentially with the length of segment studied. This suggests that a small polarity of auxin transport occurs in each cell, and that when auxin is being transported along an organ through a series of cells there is an amplification of the

individual small polarities of transport. It has been calculated for *Zea mays* coleoptile segments that the observed overall polarity of auxin transport could result from each cell in the longitudinal series possessing a 1 to 10 per cent basipetal polarity (i.e. a single cell need secrete no more than 1.01 to 1.10 times as much auxin basally than apically).

The mechanism by which polar transport of auxin occurs is not yet understood. For many years it has been known that the polar transport of auxin involves metabolic processes because, (a) it takes place more rapidly than can be explained by simple physical diffusion, (b) velocities of auxin movement are greatest at temperatures favourable for the activities of most enzymes (20–30°C), (c) respiration, especially aerobic respiration, is required for the maintenance of polar transport, (d) auxin can be transported in a polar manner against its own concentration gradient, and (e) the polar transport system appears to be specific for molecules that possess biological activity similar to that of IAA.

Until very recently, it has been generally considered that auxin is taken up passively by each cell, moves in the cytoplasm (and perhaps also through the vacuole) and is secreted across the plasma membrane at one end by a *carrier-mediated, energy-requiring* mechanism. In other words, that a localized *active-transport* system is responsible for the secretion of auxin and hence its polar transport.

An alternative theory of polar auxin transport has recently been suggested. Since cells are more permeable to undissociated auxin molecules (IAAH) than to auxin anions (IAA⁻), they will accumulate auxin when the pH of the cytoplasm is above that of the walls. Once inside the cell the auxin dissociates into IAA⁻. Polarity will occur if the ratio of the permeability of the anion to the undissociated auxin, i.e. $P_{\text{IAAH}}/P_{\text{IAA}^-}$ (P = permeability coefficient) is relatively greater at one end of the cell than the other. Moreover, this hypothesis assumes that not only is uptake of IAAH a "passive" diffusion process, but also that *efflux* of IAA⁻ occurs in response to the existence of gradients in pH and electrical potential across the plasma membrane. Such transmembrane gradients do exist and can be measured, and so long as the pH and/or the (negative) electrical potential in the cytoplasm is maintained at a higher level than in the free space outside, then diffusion of IAAH along its concentration gradient into the cell will be balanced by the efflux of IAA⁻ down an outwardly directed electrochemical potential gradient. Thus, whereas carrier-mediated secretion requires expenditure of energy to transport auxin against a concentration gradient, the "new" theory postulates that the movement of auxin is thermodynamically "downhill" and that metabolic energy will only be required to maintain pH and electrical gradients and polar permeability. Although some experimental evidence tends to support the "new" hypothesis other evidence is consistent with the carrier-mediated hypothesis and hence more investigation will be required to resolve this problem.

The Transport of Gibberellins in Plants

Far fewer studies have been made of gibberellin translocation than of auxin transport, but these have nevertheless provided fairly convincing evidence that gibberellin transport

is not polar in nature, except possibly in leaf petioles. Thus, it is generally noticed that gibberellins applied to any one part of a plant elicit developmental responses in all other regions of shoot and root, which provides indirect evidence for non-polar transport of the hormone. More direct evidence that gibberellins are able to move freely in all directions within the plant has been obtained by use of radioactive gibberellins (Fig. 5.2).

Experimentation on the transport of gibberellins has not usually involved use of the agar "donor" and "receiver" blocks technique that has provided such a convenient and major

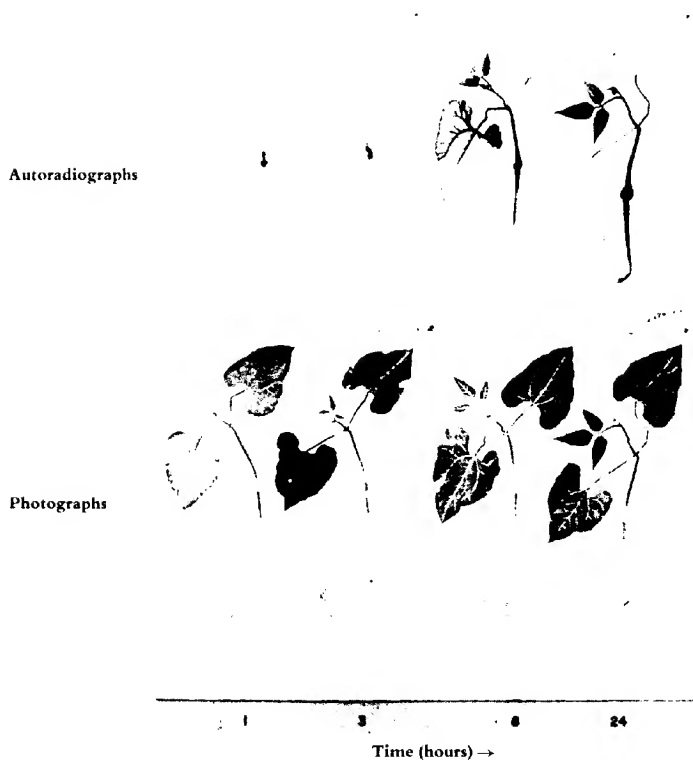


FIG. 5.2. Apparently free, non-polar, transport of gibberellin in bean plants. ^{14}C -labelled gibberellin was applied to the cotyledonary node of the plant, and its distribution with time within the plant followed by autoradiography. (From G. Zweig *et al.*, *Adv. Chem. Ser.* 28, 122-34, 1961. Original print donated by Dr. Gunter Zweig.)

tool in studies of auxin transport in segments of stems, coleoptiles, etc. The principal reason for this is that it has been found that ^{14}C - or ^3H -labelled gibberellins are taken up readily enough by tissue segments, but are not exported into agar receiver blocks to any significant extent. The reasons for this are not known, but it provides an interesting contrast to the facility with which ^{14}C -auxins pass out into such receiver blocks, in that it emphasizes the secretory nature of polar transport of auxins.

It is considered that the movement of gibberellins within the plant occurs through the normal general circulatory system of the phloem and xylem vascular tissues, since they have been detected in both xylem and phloem sap. However, one aspect of the transport of endogenous gibberellins for which phloem or xylem transport does not readily account, is the movement of gibberellins from their putative major regions of synthesis in young growing leaves downwards into and along the stem. Such young leaves are net importers of organic and inorganic materials, and the direction of flow of solutes in phloem and xylem is acropetal (upwards) in the apical part of the shoot. If phloem transport occurs in response to demand for assimilates by "sinks", then it is impossible to visualize basipetal gibberellin transport in the phloem of the apical region of the shoot.

Cytokinin Translocation

No clear picture can be presented of the transport of cytokinins in plants. Available relevant evidence is slight, fragmentary and often contradictory.

Experiments that have demonstrated the role of roots in maintaining a supply of cytokinins to leaves and preventing their premature senescence (Chapter 12) are clearly indicative of an upward transport of cytokinins in the stem. Furthermore, cytokinins are known to be present in the xylem sap ascending from the root system. On the other hand, cytokinins that are synthesized in young developing fruits do not appear to be transported out at all. Similarly, numerous studies with exogenous cytokinins such as kinetin have indicated that they may remain for some considerable time in the localized region of application, even though general metabolite transport may be taking place away from the point of application of the cytokinin./

A certain amount of evidence suggests that cytokinins may be transported not as free purines, but in conjugated forms, such as ribosides or glucosides, both of which have been shown to be present in xylem and phloem sap.

Ethylene Transport

Although as a substance of low molecular weight it might be expected that ethylene would move freely through plant tissues by normal physical diffusion, this does not appear to be the case. Thus, in broad bean for example, the "resistance" to longitudinal movement

up or down the plant is such that lateral emanation effectively isolates different parts of the plant from one another. Similarly if a leaf is fed with ethylene, only a small proportion of the gas which passes into the leaf ever reaches the stem, the remainder being emanated from the petiole. Hence, ethylene does not move between different parts of the plant in physiologically significant amounts.

Despite the fact that ethylene is not translocated to any significant extent, nevertheless it appears that changes in ethylene levels in one part of a plant can influence those in another. Thus an increase in ethylene levels in the roots can induce increased levels in the shoot apex. The mechanism of this effect is not understood at present.

Transport of Abscissic Acid

Application of exogenous ABA to mature leaves, or even to roots, can result in developmental responses such as growth inhibition in all other parts, e.g. the shoot apex or the vascular cambium. Such observations indicate that, like gibberellins, ABA is able to move freely in all directions within the plant. Also, studies with ^{14}C -labelled ABA have given no clear evidence of polarity of ABA transport in stem and coleoptile segments, but have provided evidence to suggest that in root segments ABA is transported preferentially in a basipetal manner (i.e. away from the root apex). Some research has indicated that ABA may be synthesized in the root cap, and a basipetal mode of transport in roots would therefore provide a mechanism for this ABA to reach and influence the elongating region of the root (p. 179).

GROWTH HORMONES AND STEM GROWTH

We shall now consider the role of growth hormones in the control of stem growth. Although effects on elongation growth are possibly the most intensively investigated aspects of hormone action in plants, it is extremely difficult to draw any sort of clear picture of the overall mechanism by which stem elongation is normally regulated. Each of the five known categories of growth hormone can certainly influence stem growth. The principal problem is to visualize how they interact, for their individual effects appear to overlap, duplicate, reinforce, or antagonize one another to a bewildering extent. Furthermore, not only are we still uncertain of the sites of synthesis of some types of growth hormone (especially the cytokinins and abscissic acid), but we have only sketchy information on the factors which determine patterns of transport of hormones other than auxin. In our present state of knowledge we are therefore compelled to consider individually the evidence for the involvement in stem elongation of each class of hormone, and to attempt only briefly the synthesis of available information for all hormones towards a general understanding of the control of stem elongation growth.

Auxin and Internode Elongation

As we considered in Chapter 3, the discovery of the natural auxin, IAA, took place as a result of experiments on phototropism and elongation growth in etiolated coleoptiles. In particular, observations that removal of the apical end of a coleoptile resulted in suppression or cessation of elongation growth in the remaining coleoptile, and that IAA could substitute for the tip, led to the conclusion that the apical part of a coleoptile normally supplies auxin to the newly formed cells and that the auxin is necessary for elongation growth of those cells. Convenient though are etiolated coleoptiles in experiments on the role of apically-synthesized auxin in the regulation of extension growth, one must remember that they are modified, tubular leaves, and that results obtained by their use should be applied only with caution to the problem of how internode elongation growth is normally controlled.

It is now generally believed that auxin which is involved in the control of internode growth is synthesized in young growing leaves (or in cotyledons in very young seedlings) from where it is translocated into and basipetally down the stem. Evidence for this view is largely circumstantial, but includes positive identification by mass-spectrometry of IAA in young leaves of bean plants, and measurements of quantities of unidentified auxin (presumably IAA) diffusing out from the petioles of variously aged leaves of *Coleus* plants (Fig. 5.4).

It has been found that those internode tissues which are most rapidly elongating contain the highest levels of diffusible endogenous auxin (Fig. 5.3A), which is consistent with the view that auxin is required for elongation growth in internodal regions.

Further evidence that auxin is concerned in the control of stem growth is afforded by experiments in which isolated segments of internodes are used. Such segments have been deprived of their supply of auxin from the apical region of the shoot, and consequently the effects of known concentrations of auxin can be measured. Thus, if the internode segments are floated in an appropriate solution of auxin, then they will grow more than when placed in water alone. Further, the amount of extension growth that occurs is proportional to the logarithm of the concentration of auxin in solution (Fig. 5.5A). It can be seen that stem segments show increased growth with increasing concentrations of IAA up to approximately 5×10^{-5} M. At concentrations greater than this less growth occurs, until at about 10^{-3} M IAA and above inhibition of excised stem segment elongation occurs, so that less growth takes place than in the control segments to which no IAA was applied. Consequently, we can say that for stem tissues there is a concentration of auxin which is *optimal* for cell elongation. Concentrations greater than this are, therefore, *supra-optimal* and lower concentrations are *sub-optimal*. In some species, such as pea, much higher concentrations of auxin are required to produce supra-optimal inhibition in light-grown (green) stem segments than in etiolated stem segments of the same species.

Since application of auxin to an intact shoot (e.g. as a spray) rarely stimulates stem elongation, it may be assumed that the stem tip normally supplies sufficient auxin to the elongating internodes to maintain an optimal auxin concentration in those tissues. The possible reasons for inhibition of growth by auxins will be discussed below (p. 113).

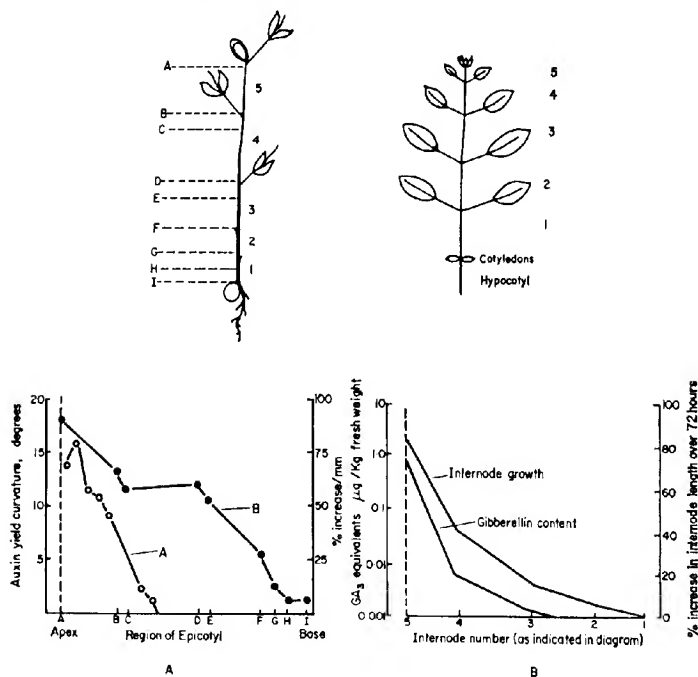


FIG. 5.3. A. The correlation between growth rate and auxin content along the length of a pea seedling stem (epicotyl). Above: diagram of a green 9-day-old "Alaska" pea seedling. The figures indicate the number of each internode, and the letters delimit regions assayed for auxin content. Below: distribution of diffusible auxin and extension growth in the epicotyl. Curve A, relative growth rates. Curve B, auxin yields. (From T. K. Scott and W. R. Briggs, *Amer. J. Bot.* **47**, 492-9, 1960).

B. The relationship between elongation rate and gibberellin content along the length of the stem of a young sunflower (*Helianthus annuus*) plant. (From R. L. Jones and I. D. J. Phillips, *Plant Physiol.* **41**, 1381, 1966.)

Gibberellins and Internode Elongation

There are several pieces of evidence which indicate that gibberellins, as well as auxins, are involved in extension growth of plant tissues. The most striking and characteristic response of a plant treated with a gibberellin such as gibberellic acid is that stem elongation is stimulated, so that the treated plant becomes taller than it would normally. This response of the stem is usually due to an increased elongation of the internodes and there is generally no increase in the number of internodes formed. The increased internode length is a consequence of increased cell extension and cell division. Thus, gibberellin treatment of intact

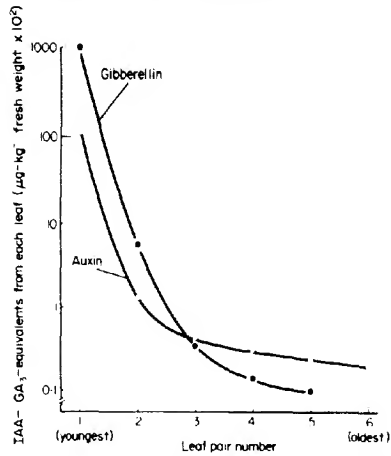


FIG. 5.4. Amounts of gibberellin and auxin transported out from leaves of various ages. Young growing leaves synthesize and export to the stem greater quantities of both types of growth hormone than do older leaves. Gibberellin data from experiments on sunflower plants (R. L. Jones and I. D. J. Phillips, *Plant Physiol.* **41**, 1381-6, 1966) and auxin data from experiments with *Coleus* plants (R. H. Wetmore and W. P. Jacobs, *Amer. J. Bot.* **40**, 272-6, 1953.)

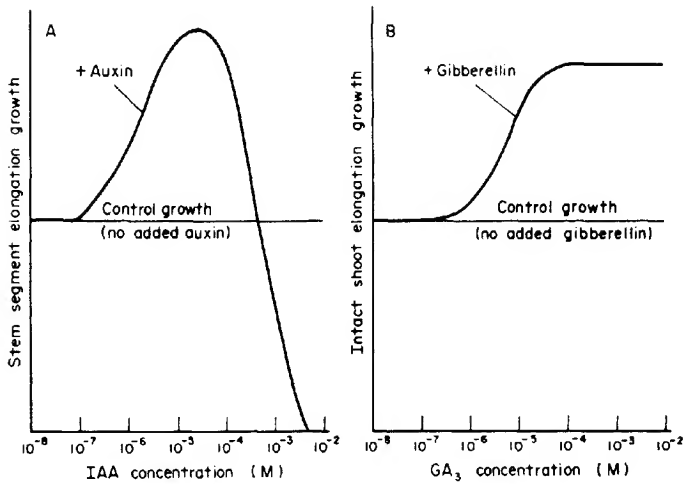


FIG. 5.5. A. Typical dose-response curve for elongation growth of stem or coleoptile segments in varying concentrations of the auxin indole-3-acetic acid (IAA). B. Typical dose-response curve for elongation of intact shoots treated with various concentrations of the gibberellin gibberellic acid (GA_3).

plants can cause enhanced elongation of existing internodal cells, and also increase the number of cells present in each internode, principally as a result of an increase in mitoses in the sub-apical region of the stem (Fig. 5.6).

The magnitude of the stem-elongation response to gibberellin varies from species to species and from variety to variety within a species. As mentioned in Chapter 3 (p. 56), the response is greatest in genetically dwarf plants—so much so, that following gibberellin treatment dwarf varieties often grow to the same height as that of related tall varieties (Fig. 3.3). Tall varieties of the same species respond only slightly or not at all. In contrast to the typical dose-response relationships between auxin and stem segment elongation (Fig. 5.5A), gibberellins rarely show supra-optimal inhibition of elongation, for even very high concentrations of exogenous gibberellic acid can elicit a maximum growth response (Fig. 5.5B). The reasons for this difference in pattern of response to varying auxin or gibberellin concentration are not fully understood, but supra-optimal auxin concentrations can induce increased release of ethylene (Fig. 5.9), particularly in dicotyledonous species, and it is likely that it is the additional ethylene which causes growth inhibition. Gibberellins, on the other hand, have been found to have very variable effects on ethylene production. In the majority of investigated situations, a small promotive effect of gibberellin on ethylene production has been found, but in others gibberellins had either no effect or even decreased ethylene levels.

The responsiveness of different varieties to applied gibberellin may perhaps be related to the amount of endogenous gibberellin present in the tissues. However, in some species, such as *Pisum sativum*, contradictory results have been obtained by different research groups in that some have found lower levels of gibberellins in dwarf than in tall pea varieties, whereas others have been unable to detect any quantitative difference. Consequently, one

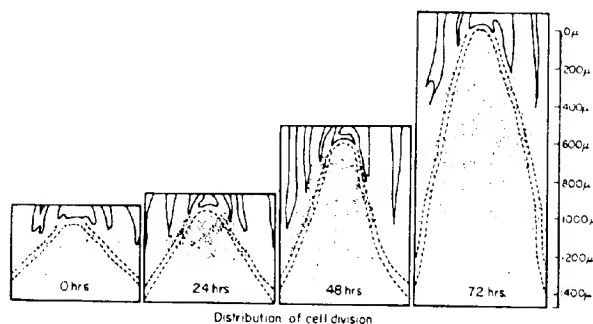


Fig. 5.6. Stimulatory effect of gibberellic acid (GA_3) upon sub-apical meristematic activity in *Samolus parviflorus*, a rosette plant. Each dot represents a sub-apical cell undergoing mitosis seen in median longitudinal sections of the stem apical region. Twenty-five micrograms of GA_3 were applied 0, 24, 48 and 72 hours previously. (Reprinted from R. M. Sachs, C. F. Bretz and A. Lang, *Amer. J. Bot.* 46, 376–84, 1959.)

cannot yet conclude that dwarfness of a plant is always due to impaired gibberellin synthesis, and further research is being conducted to resolve this question.

It is possible to extract gibberellins from various organs, and to compare the amounts present in them; also, it is possible to collect gibberellins from organs by the agar diffusion technique previously used in studies of auxins. Such studies have revealed that the apical bud of the shoot, and young leaves, synthesize and export gibberellins to the stem (Fig. 5.4). Also, a positive correlation has been obtained in sunflower (*Helianthus annuus*) between the growth rates of internodes of different ages and the gibberellin contents of the same internodes (Fig. 5.3B). Thus, as with auxins, endogenous gibberellins are present in highest concentration in those regions of the stem which are undergoing most rapid extension growth, providing strong circumstantial evidence that gibberellins are concerned in the normal control of stem extension growth. Cytokinins may be involved in the control of cell division rates at the stem apex, but we have no evidence that internode extension is in any way under a direct influence of this class of hormone.

Auxin-Gibberellin Interactions in Stem Elongation

Following the discovery of gibberellins and the realization that these hormones occur naturally in higher plants (see Chapter 3), many plant physiologists were led to study the interactions of gibberellins and auxins in stem and coleoptile extension growth.

Spraying an intact plant with gibberellin was found to enhance internode extension growth, whereas it was already known that similar treatment with an auxin rarely induced greater internode elongation. Conversely, if young internodes or coleoptile segments were excised and floated in solution, then it was noted that the opposite usually occurred; auxins stimulated internode or coleoptile segment elongation but gibberellins had little or no effect. However, when gibberellin and auxin were present in solution together, then the elongation of excised segments was much greater than when auxin alone was supplied (Fig. 5.7). In other words, for the characteristic effects of gibberellins on stem elongation to appear, auxin also has to be present. The growth-promoting action of gibberellins when applied to intact plants would, therefore, be a consequence of an interaction between supplied exogenous gibberellin and the natural endogenous auxins present. Because of such observations, it was proposed that gibberellins exert their physiological effects through some "auxin-mediated" mechanism. Thus, there is good evidence that application of gibberellins leads to increased endogenous auxin levels, by effects on either the biosynthesis or the destruction of auxin.

However, it is now clear that gibberellins are a class of growth hormone in their own right. If gibberellin effects were due solely to an influence on auxin activity then it would be expected that these effects would always be the same as those produced by auxins. Whilst gibberellins often are able to duplicate the known effects of auxins (e.g. induction of parthenocarpic fruits, promotion of cambial activity), there are many other examples of gibberellins having physiological effects not possessed by auxins (e.g. promotion of stem

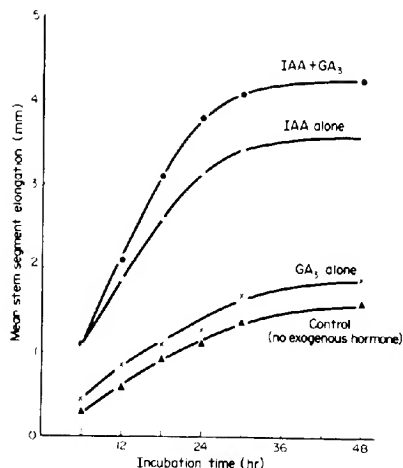


FIG. 5.7. Effects of auxin ($10 \mu\text{g ml}^{-1}$ IAA) and gibberellin ($10 \mu\text{g ml}^{-1}$ GA₃) on elongation growth in excised pea internode segments. Elongation of stem segments is much more markedly enhanced by auxin than by gibberellin, but the addition of both auxin and gibberellin results in more growth than with auxin alone. The interaction between exogenous auxin and gibberellin is sometimes additive and sometimes synergistic. (Adapted from P. W. Brian and H. G. Hemming, *Ann. Bot.*, n.s., **22**, 1-17, 1958.)

elongation in intact plants, breaking dormancy of buds or seeds, stimulation of mesophyll growth), or sometimes instances where gibberellins have the opposite effects of auxins (e.g. auxins promote but gibberellins inhibit root initiation in stem cuttings).

Nevertheless, it is apparent that interactions between auxins and gibberellins occur in many physiological responses apart from extension growth. For example, auxin and gibberellin together are often more effective in inducing the development of parthenocarpic fruits and in stimulating cambial activity (p. 120) than is either type of hormone on its own.

Ethylene and Internode Elongation

Impressive evidence has accumulated in recent years which suggests that ethylene may play some part in the control of stem growth. With most species, exposure of stems or isolated internodes to ethylene reduces cell elongation (Fig. 5.8) but enhances isodiametric cell expansion. Such ethylene-treated internodes are shorter and thicker than untreated internodes. These effects of ethylene are similar to those which can be induced by high

(supra-optimal) concentrations of auxins. It is possible that auxins are not themselves inhibitors of stem elongation, but rather that at high concentrations they stimulate the synthesis of ethylene in plant tissues, which in turn, suppresses cell elongation (Fig. 5.9).

The inhibiting effect of ethylene on stem elongation is less in the presence of light than in darkness. The reasons for this are not understood, but ethylene release can be affected, probably indirectly, by red and far-red light through a phytochrome-mediated mechanism. Thus, the results of experiments done on the effects of exogenous ethylene are likely to be influenced by the light régime, in that the response to a given concentration of exogenous ethylene will depend upon the prevailing endogenous ethylene level in the tissues.

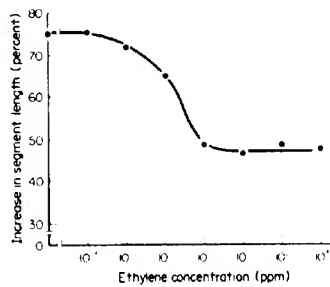


FIG. 5.8. Inhibition by ethylene of elongation growth in excised pea-stem segments. (Adapted from S. P. Burg, *Régulateurs Naturels de la Croissance Végétale*, Edition de la Rech. Sci., 1964, pp. 718-24.)

A special case of stem elongation which involves regulation by ethylene is seen in examples of those seedlings in which the terminal part of the shoot axis is hook-shaped (Fig. 8.5). The hook is presumed to aid penetration through the soil of the delicate apical tissues of the shoot. The shape of the hook is a resultant of more rapid elongation growth on the outer convex side than on the inner concave side. Exposure of seedlings to red light causes the hook to open by an equalization of growth rates on the two sides of the stem, and far-red light reverses the effect of red light. Treating red light-grown seedlings with either auxin or ethylene results in closure of the hook by inhibition of the inner side stem tissues. It has been found that ethylene acts in an intermediary capacity in both light- and auxin-controlled hook opening and closing. That is, both far-red light and auxin induce ethylene release in the apical part of the shoot, and the ethylene inhibits extension in the inner side of the hook. Red light reduces ethylene release and causes opening of the hook. However, the mechanism of hook opening and closing appears to be more complex than this, and is not yet fully understood. For example, it is known that gibberellins can also influence hook formation and that the effects of light cannot be completely explained in terms of effects on ethylene production.

In some plants, mainly species that grow well under water (e.g. rice and *Callitriche*),

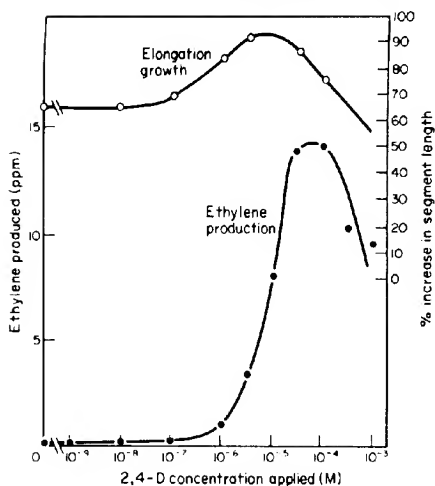


FIG. 5.9. Effects of various concentrations of the auxin 2,4-dichlorophenoxyacetic acid (2,4-D) on elongation growth (top) and ethylene synthesis (bottom) in excised segments of soybean hypocotyls. Note that maximum elongation occurred with approximately 10^{-5} M 2,4-D, but that at higher concentrations of 2,4-D the rate of ethylene production rose markedly and elongation growth decreased. It is likely that growth inhibitions in dicotyledonous species by high concentrations of auxins occur through the inhibitory effects of the auxin-induced ethylene, but this may not be so for monocotyledonous plants. (Adapted from R. E. Holm and F. B. Abeles, *Planta*, 78, 293, 1967.)

rather than being inhibited, internode and root extension occurs more rapidly in the presence of ethylene. The elongation responses to ethylene shown by these aquatic plants may have evolved as an adaptation to the very much lower rates of diffusion of ethylene in water than in air. Thus, movement of emanated ethylene away from the plant surface is considerably slower in an aquatic environment, which could result in a high ethylene concentration within submerged plant tissues. Changes in sensitivity and response to ethylene may, therefore, have occurred during evolution of aquatic plants to cope with this problem.

It is clear that ethylene interacts with the other natural growth hormones in the regulation of stem elongation growth, but much research remains to be done to unravel what is clearly a complex and sensitive control mechanism.

HORMONES AND ROOT GROWTH

It is probably impossible to think of an aspect of plant physiology which reveals greater ignorance and confusion than that which is concerned with natural regulation of growth

and differentiation in roots. By simple analogy with the situation as we understand it in coleoptiles and stems, one might expect that auxins and gibberellins are synthesized in the apical part of the root, and that these hormones are translocated basipetally to the cell elongation zone a few millimetres behind and there regulate growth and differentiation. However, no unequivocal evidence exists that can allow us to formulate such a view. Firstly, it is very uncertain what effects are exerted by the root tip on root elongation growth, for excision of the root apical meristem does not necessarily prevent elongation of the newly formed cells in the root. It has on occasion even been found that root-tip removal results in temporarily *stimulated* elongation of the young root cells. Secondly, although experimental evidence shows that auxin (IAA) is present in roots, its concentration is extremely low and it cannot be assumed that IAA is synthesized in root tissues. Thirdly, as we have already seen (p. 102), studies of radioactive IAA transport in roots have yielded contradictory and confusing results. Those that have demonstrated acropetal polarity of auxin transport (e.g. Fig. 5.1B) would appear to exclude the possibility of regulation of elongation growth through the action of auxin synthesized in the root apical region. There is somewhat better evidence that gibberellins may be synthesized in root tips and transported basipetally into the cell elongation zone, but even here the data are less than completely convincing. Furthermore, applications of gibberellins to plants generally have little obvious effect upon the growth of roots. Exogenous ethylene inhibits root elongation as effectively as stem elongation (except in rice, where both stem and root elongation are promoted by applied ethylene) by what appears to be an identical mechanism. Roots also synthesize ethylene, and it is possible that supra-optimal auxin concentrations inhibit root elongation through enhancement of ethylene biosynthesis in the root tissues.

Current research on geotropism in roots (see Chapter 7) is tending to emphasize the importance of the root cap as an organ of both gravity perception and the site of synthesis of growth inhibitors, including abscisic acid. Abscisic acid is transported basipetally in roots, and it is clear that the possible involvement of ABA, and perhaps other as yet unidentified growth inhibitors, will have to be considered in future work on the problem of extension growth in root cells.

Our knowledge of cytokinins and their possible functions in root growth is extremely limited. Treatment of excised roots growing in culture with a cytokinin, in combination with an auxin, results in a stimulated cell division rate. However, this usually does not lead to an increased rate of root elongation, but to a stimulation of division of cells which are destined to differentiate into vascular tissues. Thus, it appears likely that endogenous cytokinins are involved in the control of vascular development in the root, but it is not possible at present to determine any function for cytokinins in root elongation growth.

HORMONES AND THE GROWTH OF LEAVES

Once a leaf primordium has been initiated at the stem apex, it starts to grow and develop by the processes of cell division, cell enlargement and differentiation (see Chapter 2). One

can reasonably assume that these processes are under the controlling influence of growth hormones, one of which would be expected to be auxin. However, it cannot be said that auxin is involved in all aspects of leaf growth. It has been found that auxins will, depending on their concentration, either stimulate or inhibit the growth of midrib and veins but have little effect on the interveinal mesophyll tissues. At the present time little is known of the hormonal control of leaf growth, other than that auxin appears to be necessary for vein growth.

It has been suggested that a growth hormone synthesized in the roots controls mesophyll growth. Thus, it is found that if young root tips are cut off as fast as they are formed in, for example, the horseradish plant (*Amoracea lapathifolia*), normal development of the interveinal tissue fails to occur (Fig. 5.10). Evidence that roots export cytokinins and gibberellins to the shoot may provide a partial explanation for failure of mesophyll growth in such root-pruning experiments.

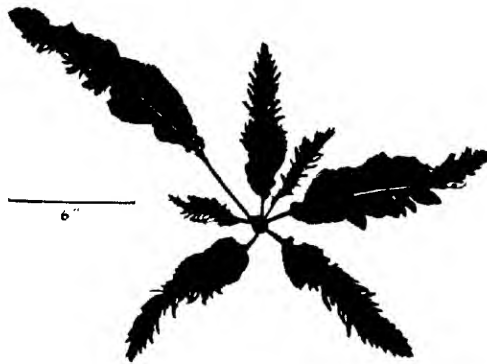


FIG. 5.10. Effect of repeated excision of root tips upon the growth of mesophyll in leaves of the horseradish plant (*Amoracea lapathifolia*). The smallest and most deeply lobed leaves are the youngest. (Photographed by Mr. J. Champion, from a plant grown by Dr. J. Dore.)

It has been found that treatment of some plant species (e.g. *Triticum* and *Phaseolus*) with gibberellic acid leads to a stimulation of leaf growth. Other plant species (e.g. *Pisum sativum*) do not respond in this way, and in fact mesophyll growth may be retarded following gibberellin treatment. In those species where gibberellin treatment does stimulate leaf growth, the mesophyll and vein tissues respond nearly equally. Similarly, excised leaves or leaf disks will, if floated on the surface of a solution of gibberellin or cytokinin, often expand due to growth of the mesophyll. Thus, generally speaking, gibberellins and cytokinins differ from auxins in their effects on leaf growth, in that the former two classes of hormone

can promote mesophyll growth whereas auxins do not. This immediately suggests that endogenous gibberellins and cytokinins are important regulators of the growth of leaves. In fact it has been found that gibberellins are normally present in leaves, and that the concentration present is closely related to the growth rate of the leaf, so that young, rapidly growing leaves contain more gibberellin than do older leaves (Fig. 5.11). Natural gibberellins may, therefore, be of importance in the control of mesophyll growth. We know little about endogenous cytokinins in leaves, although some experiments have shown that synthetic cytokinins can replace the need for a root system for healthy leaf growth.

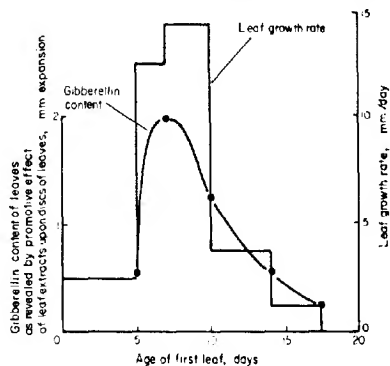


FIG. 5.11. Relationship between growth rate (histogram) and gibberellin content (graph) in first leaf of dwarf French bean (*Phaseolus vulgaris*). Maximum gibberellin content of the leaves occurred at 7 days of age, coinciding with the period of most vigorous leaf expansion. (From A. W. Wheeler, *J. Exp. Bot.* **11**, 217-26, 1960.)

In cereal and grass species, a late stage of leaf development involves unrolling of the lamina—i.e. much of longitudinal and lateral expansion of the leaf takes place with the leaf rolled into a cylindrical form with the adaxial surface innermost. The physiological basis of this leaf-unrolling process, which is under phytochrome control, is considered in more detail in Chapter 8.

We have already mentioned that certain phytochrome-controlled responses may be mediated through changes in ethylene release, and in view of the inhibitory effects which can be exerted by exogenous ethylene on leaf expansion one cannot exclude the possibility that this hormone, also, may normally participate in leaf growth and differentiation, although it should be recognized that very little evidence is available to suggest this.

The growth of leaves may, therefore, be under the controlling influence of auxins, gibberellins, cytokinins and ethylene, together with nutrients and perhaps other unknown hormonal factors, but it should be borne in mind that current knowledge is meagre and fragmentary.

HORMONES AND THE INITIATION OF VASCULAR TISSUE

We saw earlier (p. 34) that developing buds and leaves exert a stimulating effect on the development of the vascular tissue in the internodes below. Moreover, experiments with chicory (*Cichorium intybus*) callus cultures showed that when a shoot bud was grafted on to the top of the callus, differentiation of vascular tissue, connected to the base of the bud, was induced (Fig. 5.12). The stimulus arising from the bud which caused the initiation of vascular tissue was shown to be capable of passing through a layer of cellophane placed between the bud and the underlying callus tissue. Subsequently, it was shown that auxins such as IAA can produce the same effects in the callus as an implanted bud. Cambial tissue, as well as xylem and phloem, can be induced in several types of callus tissue by the application of IAA, especially if applied in association with sucrose. Moreover, extended application of IAA to stems and roots of pea will induce the formation of vascular tissue in the cortex (Fig. 5.13). These observations suggest that the stimulating effect of developing leaves on the initiation of vascular tissue may be mediated partly by the endogenous IAA produced by the leaves.

This conclusion is also supported by observations on the regeneration of vascular tissue following wounding. If one of the vascular bundles of *Colens* is severed by a lateral cut, regeneration of the strand occurs through the pith (and connects the severed upper and lower ends of the bundles (Fig. 5.14)). The pattern of regeneration is basipetal, i.e. it starts from the upper severed end of the original vascular bundle and travels from cell to cell diagonally through the pith. The regenerated strand is formed by the differentiation of parenchymatous pith cells into xylem and phloem, but subsequently cell division occurs and a cambium is formed. It has been shown that the presence of a leaf above the cut in the stem is essential for the regeneration of the strand, but the effect of a leaf can be replaced by

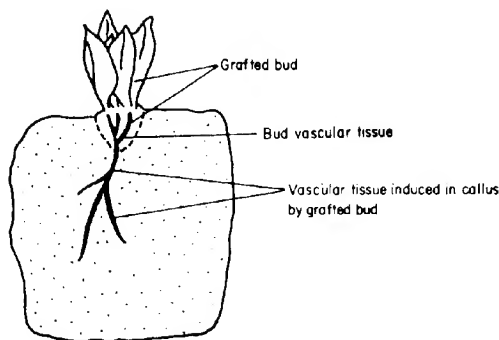


FIG. 5.12. The induction of vascular tissue in chicory (*Cichorium intybus*) callus by a chicory shoot bud implanted on the upper surface. The regenerated vascular tissue in the callus becomes connected with that of the bud itself. (After G. Camus, *Compt. Rend. Acad. Sci. Paris*, **219**, 34, 1944.)

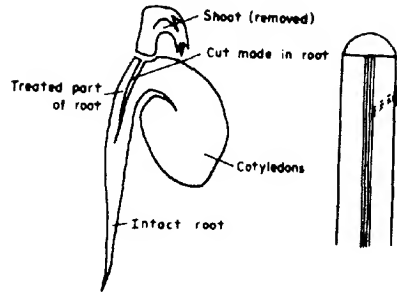


FIG. 5.13. Induction of xylem by auxin in pea roots. *Left*: 3-day-old seedling illustrating the cuts made to separate the part of the root used for experiments. *Right*: treated roots in face view. IAA applied laterally at the position indicated by black protruberance. The dotted lines indicate newly formed xylem through the cortex. (Adapted from T. Sachs, *Ann. Bot.* **32**, 781-90, 1968.)

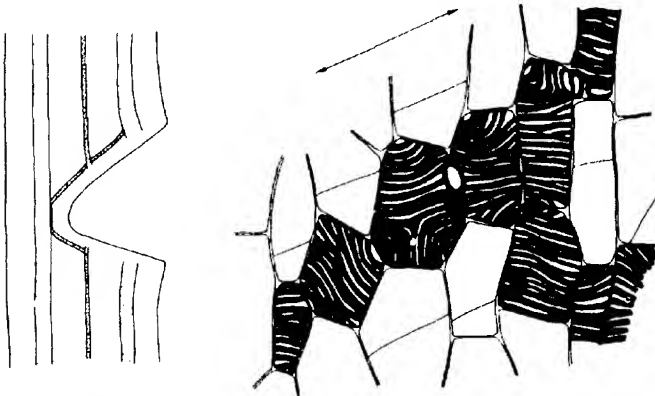


FIG. 5.14. Vascular regeneration in *Coleus*. *Left*: regeneration of connection between several vascular bundles in stem. *Right*: differentiation of parenchyma cells into reticulate xylem cells in the development of this strand. Arrow shows the direction of its development. (From E. W. Sinnott and R. Bloch, *Amer. J. Bot.* **32**, 151, 1945.)

applying auxin to the petiole stump, suggesting that auxin stimulates the differentiation of xylem and phloem and the formation of a cambial layer in the pith cells.

The xylem elements are formed by the development of reticulate lignified thickenings on the walls of the pith cells. During the early stages of differentiation of these cells granular strands (shown to contain microtubules which lie parallel to the future thickenings in the wall) appear in the surface layers of the cytoplasm and mark out the position of the lignified

bands which later develop on the walls. Moreover, the thickenings in one cell are opposite those of the next, so that the pattern can be observed to extend across the boundary wall between two adjacent cells. These observations strongly suggest that the pattern of differentiation in one cell may induce similar changes in an adjacent cell, a phenomenon known as *homeogenetic induction*. How this homeogenetic induction is achieved is not known, but it may involve other factors, in addition to IAA.

HORMONES AND CAMBIAL ACTIVITY

As well as stimulating the initiation of vascular tissue, hormones also appear to play an important role in cambial activity. The first demonstration that auxin stimulates the cambium to divide came from experiments performed by Snow in 1935 with sunflower (*Helianthus annuus*) plants which had been "decapitated" (i.e. the apical bud was excised). In such decapitated plants, it was found that the fascicular cambium of young internodes failed to divide, and also that interfascicular cambium did not form. In other words, excision of the apical bud prevented the formation of secondary vascular tissues (xylem and phloem). However, when IAA was applied to the upper cut end of the stem in decapitated plants, then normal cambial activity and secondary thickening resulted, suggesting that both the initiation of cambium in the stem, and its activity once formed, depend upon a supply of auxin from the apical bud.

On the other hand, a number of investigators have obtained results that suggest that the initiation and activity of the cambium may not always be regulated by auxin from young leaves. For example, a series of experiments by Siebers during the early 1970 showed that the formation of interfascicular cambium still occurred in decapitated young hypocotyls of *Ricinus communis*, or even in isolated segments of hypocotyl tissue, and he concluded that any requirement for growth hormones in cambium *initiation* must be satisfied by sources within the hypocotyl itself. However, Siebers did find that addition of IAA, GA₃, or kinetin resulted in enhanced development of the cambium once it had formed, which indicated that these types of growth hormone may normally be concerned in the regulation of cambial *activity* in the hypocotyl. Of these growth hormones, GA₃ was found to have by far the greatest stimulatory effects on cambium development and activity, and sugar, in the form of sucrose, was also important. These findings, and similar ones by other workers, suggest that one cannot regard the apical bud as the only source of auxin in the regulation of cambial initiation and activity in stems of herbaceous plants.

Rather more is known of the regulatory factors concerned in cambial activity in woody species of temperate zones. In these, there normally occurs seasonal variation in the rate of cell division activity in the vascular cambium of both shoot and root, and differences in the patterns of differentiation of cambial derivatives at different times of the year. During the winter months there is no cambial activity in such trees, but in the spring cell-division activity starts again and the newly formed cells differentiate into xylem and phloem elements.

In diffuse-porous angiospermous trees (in which all xylem vessels have a similar diameter no matter when during the growing season they develop) such as sycamore (*Acer pseudoplatanus*), cambial activity in the spring commences immediately below expanding buds and then spreads slowly *downwards*, i.e. basipetally, through the twigs to the branches, trunk and eventually into the roots. An acropetal wave of cambial activity occurs only in the roots. Thus, the initiation of cambial activity follows the same pattern as polar movement of auxin in stems and roots. Young growing buds in spring are known to transmit relatively large amounts of auxin to the stem tissues, and it therefore appears likely that the reawakening of cambial activity in diffuse porous species is a response to auxin from the buds. This idea is supported by various pieces of evidence. Thus, disbudding prevents the onset of cambial activity, but application of auxin to the upper end of a disbudded twig results in normal basipetal spread of cambial activity.

Not only is the activation of cambium regulated by auxin, but the differentiation of its derivatives is also affected by prevailing levels of auxin. It is also known that auxin is not the only hormonal regulator of cambial activity and vascular tissue differentiation. This is most easily and convincingly demonstrated in experiments in which lengths of stem from a diffuse-porous species are taken in early spring, before bud expansion has started, removing the buds, and applying growth hormones, either dispersed in lanolin or in aqueous solution, to the upper cut ends of the stem segments. After about 2 weeks the stems are sectioned for observations on cambial activity. With no applied hormone there is no cambial division, but if IAA is applied there is limited cambial division and some differentiation of new xylem elements can be observed (Fig. 5.15). If GA_3 alone is applied, cambial division occurs, but the derivative cells on the inner (xylem) side remain undifferentiated and retain their protoplasmic contents; however, careful observation shows that some new phloem, with differentiated sieve tubes, is formed in response to GA_3 . When IAA and GA_3 are applied together, there is greatly increased cambial division and normal, differentiated xylem and phloem are formed. By measuring the width of the new xylem and phloem it is possible to study the interaction of auxin and gibberellin, and other regulators, in a quantitative manner (Fig. 5.16). Experiments such as these suggest that not only can the rate of cell division in the cambium be regulated by levels of auxin and gibberellin, but also that the production and differentiation of xylem initials is favoured by a relatively high ratio of auxin to gibberellin, whereas phloem formation occurs when the gibberellin level is high.

This latter conclusion might be thought to indicate that the differentiation of cambial derivative cells into xylem on the inner side and into phloem on the outer side is determined by differences in the ratios of auxin to gibberellin concentrations on the two sides. However, a different conclusion is indicated by the results of experiments by Siebers, who carried out experiments with young seedlings of *Ricinus communis*. In this species, as in many other herbaceous plants, cambial zones are formed in the ground tissue ("interfascicular tissue") separating the primary vascular bundles and are connected to the cambia of these bundles, so forming a continuous cambial ring.

Siebers cut out small pieces of interfascicular tissue from young hypocotyls at a stage before there was any sign of the formation of interfascicular cambium. These pieces were



FIG. 5.15. Effect of auxin and gibberellin on cambial activity in poplar (*Populus robusta*) twigs. Top: twig treated with gibberellic acid (GA_3); centre: treated with indole-3-acetic acid (IAA); below: treated with IAA and GA_3 in combination.

reversed and replaced in the hypocotyls. Later examination revealed that interfascicular cambium was initiated in the reversed pieces of tissue, but the pattern of differentiation had been reversed so that xylem formed on the *outside* and phloem on the *inside*. Moreover, no direct connection was established between the interfascicular cambium and that of the primary vascular bundles. These observations suggest that although the originally complete ring of procambium at the shoot apex (p. 38) breaks down into separate strands (each of

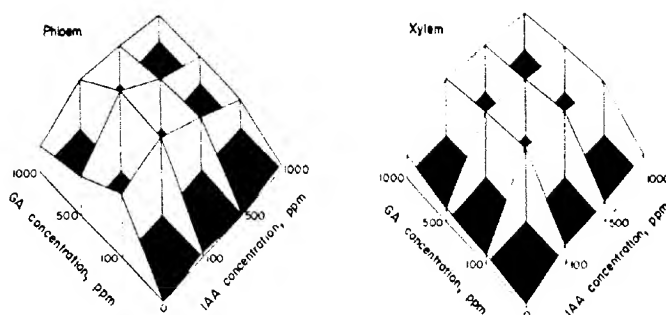


FIG. 5.16. Quantitative effects of IAA and GA_3 on cambial activity of poplar. The two hormones were applied in lanolin in various combinations at the concentrations shown. Vertical axis indicates width of new xylem and phloem in eye-piece units. See text for explanation. (From P. F. Wareing, C. Hanney and J. Digby, in *The Formation of Wood in Forest Trees*, Academic Press, New York, pp. 323-44, 1964.)

which develops into a primary vascular bundle), the regions between strands retain some propensity to develop later into cambium, even though the cells there may be morphologically indistinguishable from those of the surrounding ground tissue. Moreover, the normal pattern of differentiation of cambial derivatives (giving xylem on the inside and phloem on the outside) appears to be determined by the potentialities of the cells themselves and not by extrinsic factors, such as hormones, although the latter, especially IAA and gibberellins, are necessary for cambial division and cell differentiation.

Little is known of the role of cytokinins in normal cambial activity, but studies with isolated pea-stem sections have indicated that these hormones can also stimulate cambial division and increase the lignification of the developing xylem cells. Ethylene and abscisic acid also affect cambial activity when applied to plants, but there is not yet any evidence that these substances play a regulatory role in the natural control of cambial division and differentiation of vascular tissue.

Cambial activity is known to be affected by various non-hormonal factors, such as sugars and water availability, as well as by environmental factors, such as temperature, a fact which forms the basis of "tree-ring analysis" for dating in archaeological studies and for the study of long-term climatic changes.

HORMONES AND FRUIT GROWTH

Probably because of the economic importance of fruits, the physiology of their growth, development and ripening has been very intensively studied. Most botany textbooks classify "fruits" according to a rigid and complicated system based upon morphological

characters, but J. P. Nitsch has pointed out that a fruit is a physiological rather than a morphological entity. To the physiologist, and incidentally the layman, a "fruit" is simply a structure which arises by development of tissues which support the ovules. Nitsch has pointed out that such a view is valid even for seedless fruits because ovules were initially present in them.

The early growth of the ovary, which occurs during development of the flower, involves cell division but little cell vacuolation. In many species, cell division ceases at or shortly after anthesis (flower opening) and the subsequent growth of the fruit following pollination is primarily due to an increase in cell size rather than in cell number. For example, in tomato (*Lycopersicon esculentum*) and blackcurrant (*Ribes nigrum*), cell division ceases at anthesis and the whole of the subsequent growth is due to cell expansion. In such species, the final size will be a function of the number of cells already present in the ovary at anthesis. On the other hand, in some species (e.g. apple) cell division may continue for a time after pollination has occurred.

Fruit cells may enlarge by vacuolation to relatively enormous sizes, and mature fruits of water melon (*Citrullus vulgaris*) consist mainly of cells so large that they are distinguishable individually by the naked eye.

Fruit Set

The early development of the ovule and ovary takes place along with other aspects of flower development (p. 43). In some species, the ovary ceases growth at the time of, or before, anthesis (Fig. 5.17). In others, growth goes on for a time after anthesis and prior to

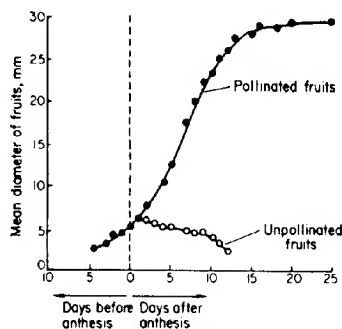


FIG. 5.17. Growth curves of ovary of *Cucumis anguria*. Growth in unpollinated ovaries ceases soon after anthesis, but pollinated ovaries show a typical sigmoid growth curve. The decrease in ovary diameter seen in unpollinated ovaries is due to "shrivelling". (From J. P. Nitsch, *Quarterly Rev. Biol.* **27**, 33-57, 1952.)

pollination. In both cases, further growth of the ovary takes place only if pollination is effected. Should, for some reason, pollination not occur, then growth and development of the fruit ceases. Failure of the pollination mechanism usually results in the shedding from the plant of the unfertilized flower, often mediated by the formation of a separation, or abscission layer (see p. 298) in the peduncle. Successful pollination, on the other hand, is followed by rapid growth of the ovary and fruit development begins (Fig. 5.17). At the same time, the petals and stamens wither and often abscind. The start of ovule growth and withering of stamens and petals marks the start of fruit development, and this phase is often called *fruit set*. The process of pollination, whether or not it is followed by fertilization, is apparently sufficient to cause an initial stimulation of growth in the ovary and other parts of the future fruit. This is shown by the fact that in many fleshy fruits the increase in ovary growth may start before there has been sufficient time for fertilization to occur. Moreover, even "foreign" pollen, derived from an unrelated species and hence unable to effect fertilization, may cause marked stimulation of ovary growth.

The stimulatory effect of pollen on ovary growth appears to be due to the auxin which it contains. In 1909, Fitting observed that water extracts of orchid pollen were able to induce swelling of unfertilized orchid ovaries and withering of the petals, and it was shown later that pollen is a rich source of auxin. Finally, it was shown that pure preparations of IAA would, if applied to unfertilized flowers of a number of species, induce fruit set in the absence of any pollen. Among the species in which fruits can be set by auxins are tomato, pepper, tobacco, holly, figs and blackberry. In these species, fruits which have been set by treating unpollinated flowers are seedless. The production of such seedless fruits is called *parthenocarp*. Recently it has been found that ethylene will simulate the effects of pollen on ovary swelling and petal withering in various species. Thus, it is possible that some effects of auxins on ovaries are mediated through influences on ethylene production.

Fruit Growth

Although pollination may stimulate the initial swelling of the fruit, in most species further development of the fruit appears to be dependent upon the presence of developing seeds, and hence can only occur when fertilization is effected. Thus, in many fruits, such as grapes, blackcurrants, tomatoes, apples and pears, strong correlations exist between the final size of the fruit and the number of fully developed seeds it contains. In the case of the strawberry, Nitsch showed by elegant surgical experiments that receptacle growth is dependent upon achene development (Fig. 5.18).

The interaction between the growth of the ovary and that of the embryo and endosperm is shown by the changing growth rates of these different parts of the fruit at different stages of development. In some instances the growth curve for the fruit is sigmoid (e.g. in apple) and in others it is doubly sigmoid (Fig. 5.19). In peach, the changes in growth rate of the pericarp can apparently be correlated with changes in growth rate of the developing seeds. The stimulatory effect of the developing seeds upon the growth of pericarp tissue appears

to be due, at least partly, to the auxin which they produce. Developing seeds are rich sources of auxin and it has been shown that there is a declining gradient of auxin concentration within the tissues of the fruit, with the highest concentration in the seeds, less in the placenta and least in the carpel wall. This gradient is consistent with the view that auxin is produced in the developing seeds and moves outwards from there to the other parts of the fruit.

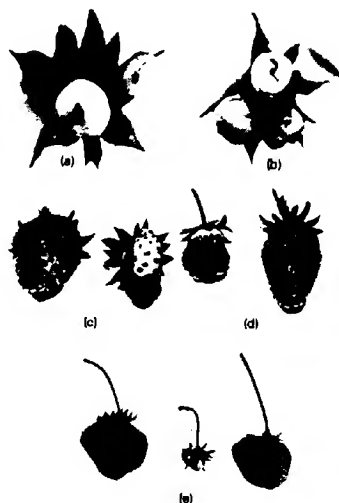


FIG. 5.18. Correlation between achene development and receptacle growth in strawberry. A: only one fertilized achene present; B: three fertilized achenes present; C: *left*, control fruit; *right*, fruit with three vertical rows of achenes. D: *left*, fruit with two rows of achenes; *right*, control fruit. E: three strawberries of the same age; *left*, control; *middle*, all achenes removed and receptacle smeared with lanolin paste; *right*, all achenes removed and receptacle smeared with lanolin paste containing 100 ppm of the auxin β -naphthoxyacetic acid. (Original prints kindly supplied by Dr. J. P. Nitsch, Laboratoire du Phytotron, Gif-sur-Yvette, France. From *Amer. J. Bot.* **37**, 211-15, 1950.)

A good example of the relationships that have been found between endogenous auxins and fruit growth has come from studies of berries of the blackcurrant, which show a double sigmoid growth curve (Fig. 5.19). Two auxins were found in the berries, one acidic (possible IAA) and the other neutral (possibly IAN). These two auxins were found to be produced mainly at the times of endosperm and embryo development, which in turn coincided with the times of maximal growth rate of the fruit. It appears likely, therefore, that the first grand period of fruit growth in blackcurrant is under the control of auxins produced in the developing endosperm, and that the second grand period is induced by auxin originating in

the growing embryo. Similar patterns of auxin production in relation to fruit growth have been reported for other species by several workers, but there are other conflicting reports that no direct correlation occurs in some species between auxin content and fruit growth rates.

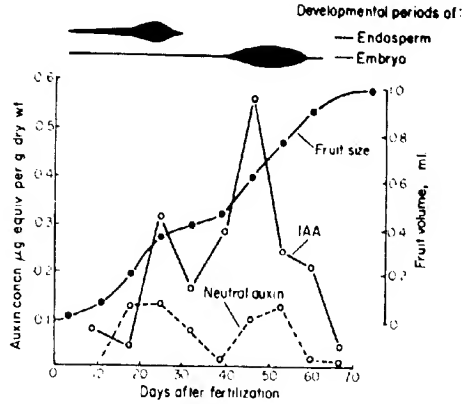


FIG. 5.19. Changes in concentration of indole-3-acetic acid (IAA) and an unidentified neutral auxin in the blackcurrant berry in relation to the double sigmoid growth curve of the berry and the main developmental periods of the endosperm and embryo. (From S. T. C. Wright, *J. Hort. Sci.* **31**, 196, 1956.)

It is important to realize that despite the evidence implicating auxins in flower and fruit growth, there is also a considerable likelihood that auxins are not the only hormones involved. It has proved impossible with many plant species to induce parthenocarpic fruit development by auxin treatment, but it has proved possible to do this by spraying the flowers with a solution of gibberellin (e.g. in members of the genus *Prunus* such as cherries, peaches and almonds). However, indicative as the effects of applied gibberellins on fruit development may be, this does not in itself provide incontrovertible evidence that internally produced gibberellins are concerned in the normal growth of fruits and seeds. Consequently, studies have been made of the gibberellin contents of various fruits and seeds at different stages of their development, and in general it has been found that young developing seeds contain relatively large amounts of gibberellins. As the seeds mature and their growth slows down there is a simultaneous fall in their gibberellin content. It appears likely that gibberellins move out from the young developing seeds in a similar manner to that suggested for auxin, and that both types of hormone are involved in the control of fruit growth.

The third type of growth hormone likely to be participating in the processes of growth in a fruit are the cytokinins. As discussed earlier (Chapter 3), cytokinins are growth hormones particularly concerned in the control of cell division, and it is, therefore, likely that the active cell division which is known to occur in young developing fruits is under the controlling influence of this type of hormone. Evidence that cytokinins are involved in fruit growth is afforded by experiments which have shown the presence of cytokinins in the young fruits of apple, banana and tomato during the stages of growth in which cell division is most rapid. Thus, we may regard fruit growth and differentiation as being under the control of several types of growth hormone, a situation which probably obtains in all phases of plant development.

Parthenocarpy

We have already seen that in some species parthenocarpic fruits may be produced by treating unpollinated flowers with auxins or gibberellins. In addition to the experimental production of such fruits, natural parthenocarpy may occur in certain species. Thus, horticultural varieties of bananas, pineapples, cucumbers, tomatoes and figs exist, in which seedless fruits are normally produced without the need for any exogenous hormone. In some species, fruits are formed without pollination, while in others pollination is necessary but fertilization does not occur; in others again, fertilization occurs but the embryos abort before the fruit matures. It is not known how the growth of these seedless fruits is controlled, but it seems possible that in some cases the maternal tissues, such as the placenta, may be capable of producing auxin in the absence of normal embryos. Thus, it has been shown that the ovaries of unopened flowers of parthenocarpic varieties of orange and grape have a higher auxin content than do those of normal seeded varieties. Moreover, it has been found that young parthenocarpic fruits of cucumber contain seed-like structures, but which lack embryos and endosperm, and it is possible that these are centres of auxin production.

The production of fully developed fruits by treatment with a single application of auxin to the flowers (p. 126) also poses a number of problems, since it is not to be expected that the auxin which is applied in this way will itself be sufficient to supply the needs of the developing fruits over several weeks. However, it has been found that pollination stimulates the production of auxin by the tissues of the ovary itself in some species, such as tobacco, and it is possible that external application of auxin may similarly trigger off the production of auxin by certain tissues of the fruit, and that once this has occurred, the production of endogenous auxin will meet the requirements for the further development of the fruit. Application of auxin has been shown to lead to growth of unfertilized ovules, which develop normal looking seed coats but which contain no embryos. It is interesting to note that in some species, such as olive, hops and maize, application of auxin will stimulate initial fruit set, but further development of the fruit does not occur without pollination. Possibly, in such species the exogenous auxin does not stimulate the production of endogenous auxin necessary for further fruit development.

GROWTH CORRELATION

The various growth processes that proceed simultaneously in a plant are not independent, but are closely linked with one another. Thus, as a stem increases in length by the activities of the apical meristem, its strength is increased by activities in the cambium leading to increased girth and rigidity of the older parts, so enabling the whole shoot to stand erect. Further, as the shoot increases in size, so does the need for water and mineral nutrients increase, and this is catered for by a nicely balanced relationship between shoot growth and root growth. As the shoot increases in bulk, the size of the root system becomes proportionately larger, which allows for the additional mineral nutritional requirements of the shoot to be met. To some extent these *growth correlations*, as they are called, are explicable in terms of the availability of nutrients, or food factors, and the competition between growing regions for these substances. Thus, shoot and root growth are related to one another, and this is probably due in part to their mutual nutritional dependence; the shoot supplies the organic material which the root is unable to manufacture for itself, and in return obtains from the root the water and mineral salts to which it does not itself have direct access. In the same way, vegetative growth is very reduced when a plant is fruiting, probably mainly as a consequence of the diversion of the available food materials into the developing seeds and fruits (see Chapter 12).

This is certainly not the whole story, however. Competition for available nutrients does not adequately explain why active growth at any one time is usually restricted to only a few of the many places in a plant where it is potentially possible. An example of this is the fact that the apical meristem of a shoot is usually in a state of active growth, whereas growth of the axillary buds below it is often strongly inhibited. This characteristic of shoot growth is known as *apical dominance* or *correlative inhibition* of buds. A related phenomenon is seen in the inhibiting effect of the main root apex upon the initiation of lateral roots in the pericycle cells. Removal of the root apex causes an increase in lateral root formation in the remainder of the root, a result analogous to the effect of removing the shoot apical bud upon the growth of axillary buds. However, the mechanism by which these phenomena occur is not necessarily the same in shoot and root.

It is known that growth hormones play important roles in the correlation of growth in different parts of the plant. It is likely that all growth correlations are in one way or another affected by patterns of hormone distribution within the plant. We have already seen several examples of correlative effects in plant growth which are mediated by growth hormones, as in the stimulation of cambial activity by auxin and gibberellins arising in the buds of woody shoots, and the stimulation of fruit growth by hormones produced by the developing seeds. We shall now consider the role of hormones in apical dominance.

Apical Dominance

The apical bud of a shoot usually grows much more vigorously than the axillary buds, despite the fact that it is apparently the least favourably situated bud in relation to the supply

of organic and inorganic nutrients from the mature leaves and the root system. There is a great deal of variability between species with respect to the degree of dominance of the apical bud over the lower axillary buds. In some species, such as tall varieties of sunflower (*Helianthus annuus*), the dominance is complete and extends over almost the whole length of the stem. In others, such as tomato, the dominance of the apical bud is weaker, and the axillary buds situated only a little way below the main shoot tip grow out, resulting in a bushy shoot system.

In many species the dominance of the shoot tip becomes weaker as the plant gets older. This is seen clearly in plants such as sycamore (*Acer pseudoplatanus*) or ash (*Fraxinus excelsior*) where the early years of growth are characterized by strong growth of the leading shoot, whereas in later years a branching habit is seen. Even in herbaceous annuals there is often a weakening of apical dominance towards the end of the growing season, and in those species where the apical meristem eventually changes to produce a terminal flower, this often coincides with a release of axillary buds from correlative inhibition.

If a shoot is decapitated, i.e. the apical bud cut off, then one or more of the lower axillary buds grow out. Usually one of the outgrowing laterals becomes dominant over the others, exerting an inhibitory influence on their growth. Where this happens it is frequently the uppermost of the axillary buds which becomes the dominant shoot. Thus, some form of signal arises in an actively growing dominant shoot bud. The effects of this signal are most obviously expressed in the correlative inhibition of lateral buds and shoots, but its influence is also seen in other morphogenetic phenomena, such as in the orientation of lateral shoots, branches, rhizomes and tubers and the pattern of their differentiation.

Some experiments conducted in the first two decades of this century yielded results which indicated that young growing leaves of the apical bud synthesized a diffusible correlative growth inhibitor that normally moved through only living cells in the plant. The discovery of auxin (indole-3-acetic acid, IAA) in the early 1930s, and the realization that the young developing leaves are the primary sources of auxin in the shoot, led to investigation of the possibility that the transmission of IAA from the apical bud constitutes the correlative signal in apical dominance phenomena. It was soon found, by Thimann and Skoog in 1934, that exogenous IAA could substitute for the apical bud in maintaining inhibition of axillary buds in bean plants. This observation has been confirmed innumerable times in succeeding years for many species (Fig. 5.20). A few exceptions were later reported, especially *Coleus*, in which it was found that exogenous auxin did not exert significant inhibition on axillary bud growth. Nevertheless, it has recently been demonstrated that even in *Coleus* the inhibitory effect of apically applied exogenous auxin upon axillary shoots is revealed when the plants are rather nutritionally deficient. There seems little doubt, therefore, that the synthesis of IAA in the apical region of the shoot, probably in the young expanding leaves, and its transport down the stem constitutes a basic component of the mechanism of correlative inhibition.

The way in which auxin causes the inhibition of axillary bud growth is not, however, fully understood at the present time. It appears paradoxical that auxin, which we have so far been considering as a promoter of growth, should cause an inhibition of growth in

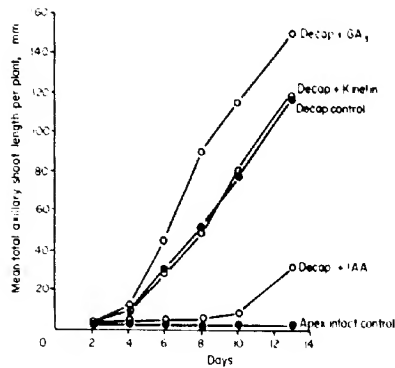


FIG. 5.20. Effect of hormones on outgrowth of axillary buds of pea plants (*Pisum sativum*) following decapitation. The hormones referred to were applied in lanolin (concentration 1000 ppm) to the decapitated stumps of the plants. The ordinates show the total growth of the three remaining axillary buds per pea plant. (L. D. J. Phillips, original data.)

axillary buds. To resolve this contradiction it was suggested by Thimann, in 1937, that the optimal auxin concentration for bud growth is lower than for stems and that the bud is inhibited by the "supra-optimal" auxin concentration (Fig. 5.5A) normally present in the stem, as a result of its synthesis in the young expanding leaves, and basipetal transport from there. This is known as the "Direct Theory" of auxin inhibition of lateral buds. It is, of course, based on the assumption that auxin enters lateral buds from the stem.

The validity of Thimann's direct theory is considered doubtful today. One of the principal objections to the theory is that determinations of the auxin contents of inhibited lateral buds in *Lupinus*, *Pisum sativum* and *Syringa vulgaris* have revealed that the auxin levels, far from being supra-optimal, are, in fact, sub-optimal. Also, it has been found that applications of low concentrations of auxin to the stumps of decapitated *Lupinus* and *Phaseolus multiflorus* plants actually accelerated growth of laterals, and that only at higher auxin concentrations did inhibition occur. Consequently, it is now generally considered that auxin does not exert its inhibitory effect on lateral buds in such a direct manner as that originally proposed by Thimann.

One of the early hypotheses for apical dominance assumed that since the apical meristem is the first-formed shoot meristem in the germinating seedling, then it would continue to command a preferential supply of metabolites as these moved along their concentration gradients. This was known as the "Nutritive Theory" of apical dominance.

If the nutritive theory is correct, one might expect that the dominance of the apical bud over the lateral buds to be most clearly manifest when a plant is deficient in nutrients, for example, when growing in soil low in necessary mineral elements. This has been clearly shown to be the case for several plants, particularly flax (*Linum usitatissimum*) in which

lateral growth is entirely suppressed by the terminal bud under conditions of mineral nutrient deficiency, whereas lateral growth occurs freely under conditions of high mineral (particularly nitrogen) nutrition. One can assume that when nutrients are freely available to the plant, there are sufficient "left over" after the apical bud has received its necessary quota to allow movement of nutrients into the lateral buds. The effect of the apical bud in flax plants growing under high or low nutrient conditions was exactly duplicated by IAA applications to the stump of decapitated plants.

Although it is undoubtedly true that the apical bud does obtain more available nutrients than the axillary buds, the simple "source-sink" explanation of the nutritive theory does not adequately explain why auxin can substitute for the apical bud in correlative inhibition of axillary buds. Further, many studies with isotopes such as ^{32}P -phosphate and ^{14}C -sucrose have demonstrated that nutrients do indeed move to, and accumulate in regions of high exogenous auxin concentration (Fig. 5.21). This "auxin-directed" transport of metabolites indicates that auxin production in the apical bud, and its basipetal transport, induces movement of available nutrients towards the region of highest auxin concentration, i.e. to the apical bud itself. It is not clear how this comes about. One suggestion has been that basipetally moving auxin in the stem inhibits the development of vascular connections (xylem and phloem) between axillary buds and main stele, so reducing the capacity of the axillary buds to obtain a supply of nutrients via the vascular system. This hypothesis therefore assumes that the lowering of the auxin content of the stem which occurs following decapitation in some way results in rapid development of bud-stem vascular connections. Quite a number of workers have tested this vascular connection hypothesis in recent years and, although results obtained are rather contradictory, the weight of accumulating evidence argues against the idea that lack of bud growth is attributable to deficiency in their vascular supply. Thus, in many cases it has been found that bud outgrowth commences within hours of decapitation of the shoot, whereas vascular connections did not develop until the buds were already growing vigorously. In other species it has been found that even completely inhibited buds appear to be served by perfectly adequate-looking xylem and phloem connections.

Other research has cast doubt upon the basic premise of the nutritive theory, in that it has been found that correlative inhibited buds contain what appear to be perfectly normal levels of major nutrients, and that feeding inhibited buds in various ways with major inorganic and organic nutrients does not induce their growth so long as the apical bud remains intact and actively growing.

However, research over the past decade has indicated that cytokinins play an important role in apical dominance. In many plants application of a cytokinin preparation directly onto a correlative inhibited bud can release that bud from inhibition in an intact plant. This suggests that inhibited buds fail to develop because they lack cytokinin. We have considered elsewhere (Chapter 12) evidence that cytokinins are synthesized mainly in the root system, but that shoot tissues require adequate supplies of these hormones for their healthy functioning. It is possible then that a role of auxin from the apical bud is to direct the transport of root synthesized cytokinins so that the apical bud receives preference over the

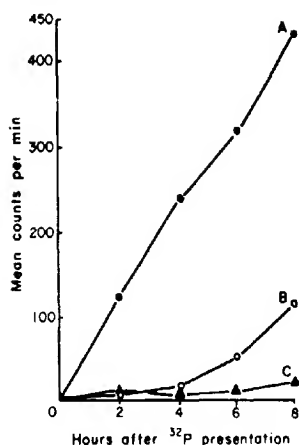


FIG. 5.21. Auxin-directed transport. The effect of IAA on the rate of accumulation of ^{32}P at the top of the stem in decapitated pea plants is shown. The radioactive ^{32}P was applied to the base of the stem just above soil level. In decapitated plants not supplied with auxin very little accumulation of ^{32}P occurred at the top of the stem (C). Application of IAA to the top of the stem immediately after excision of the apical bud (A) greatly enhanced ^{32}P accumulation. Curve B shows the effect of applying the auxin 6 hours after decapitation. (Reprinted from C. R. Davies and P. F. Wareing, *Planta*, **65**, 139–56, 1965.)

axillary buds. As considered above, correlatively inhibited buds contain low concentrations of endogenous auxin, and it is possible that it is their deficiency in auxin which prevents them obtaining adequate supplies of cytokinin from the roots. In support of this view is the fact that buds released from dominance by direct application to them of a cytokinin normally grow for only a short time, after which apical dominance is reimposed, and continued growth occurs only when auxin (and occasionally gibberellin as well it has been found) is subsequently applied to the bud. A possible explanation for this feature is that the applied exogenous cytokinin is catabolized by the growing bud before its young leaves have acquired the capacity to synthesize sufficient auxin to induce cytokinin transport into the bud. Additional evidence that cytokinin levels limit growth in correlatively inhibited buds is supplied by cytological studies that have revealed that cell division activity is suppressed in the apical meristems of such buds. Much evidence exists that cytokinins play particularly important roles in the natural regulation of cell division activity in plant cells (Chapter 3), and application of a cytokinin to an inhibited bud does indeed result in the immediate onset there of cell division activity.

Available evidence suggests that gibberellins may not be directly involved in the control of apical dominance phenomena in the way that auxins, and cytokinins are. Thus, gibberellins do not substitute for the apical bud in the correlative inhibition of axillary buds (Fig.

5.20), nor do they cause the release of buds from correlative inhibition (cytokinins can do so). Application of exogenous gibberellin to an intact plant often results in an increase in apical dominance, but this is probably due to effects on internal auxin level and distribution in the gibberellin-treated plants.

Hormonal Interaction in Stolon Development

A striking example of the importance of hormonal interaction in the control of shoot growth is seen in stolon development in the potato plant. The stolons are axillary shoots which normally arise from the basal nodes of the stem below the soil surface. They differ from the erect, leafy axillary shoots which arise on the aerial part of the shoot in that they have (1) only rudimentary leaves, (2) elongated internodes, and (3) poor development of chlorophyll (even in the light), and (4) they grow horizontally. Apical dominance evidently plays an important role in stolon development, since if the aerial shoot is decapitated and all axillary shoots are removed from it, the stolons will turn up and produce normal-looking, erect leafy shoots.

Stolons are not normally formed on the upper, leafy part of the aerial shoot, but they may be induced to develop by decapitating the shoot and applying exogenous hormones. If IAA alone is applied to the stump of a decapitated shoot, the uppermost axillary shoot is partially inhibited, whereas if GA_3 alone is applied, extension of this axillary is promoted. By contrast, when IAA and GA_3 are applied together to the decapitated stem, the development of the uppermost axillary is dramatically changed and it becomes a horizontal, stolon-like structure (Fig. 5.22). If kinetin alone is applied to the decapitated stem there is no observable effect, but if kinetin or benzyladenine is applied directly to the *tip* of a natural or of an experimentally promoted stolon, the latter very rapidly changes its pattern of development and turns upwards to become a leafy shoot (Fig. 5.22). Thus, the development of an axillary shoot of potato can be controlled very precisely by manipulating the levels of auxin, gibberellin and cytokinin. It seems likely that the natural control of stolon development involves a similar interaction between endogenous hormones.

From the various examples we have considered in this and the preceding chapter, it is evident that the correlated and integrated character of the plant is at least partially attributable to the presence of specific amounts of growth hormones at specific places and times. There remains, however, the problem of what it is which controls the precise production and distribution of growth hormones within the plant body.

PLANT GROWTH HORMONES IN AGRICULTURE AND HORTICULTURE

The fact that the mechanism of plant growth-hormone action is not understood has not prevented some of these substances becoming useful tools in agricultural and horticultural

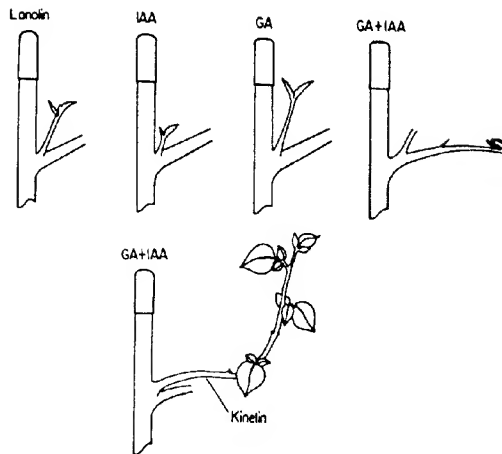


FIG. 5.22. Interaction between hormones in the control of stolon development in the wild potato species, *Solanum andigena*. Leafy aerial shoots were decapitated and hormones were applied in lanolin to the decapitated stems, as shown. The uppermost axillary shoot showed the following responses to hormone: control shoots (treated with plain lanolin) showed outgrowth of leafy axillary shoots; IAA alone caused some inhibition of growth of axillary; GA_3 alone caused some internode extension in the leafy axillary; IAA + GA_3 caused the axillary to grow as a horizontal, stolon-like shoot; when stolons were first stimulated to develop by application of IAA and GA_3 , and then kinetin was applied to the stolon tip, the axillary shoot turned upwards and showed normal leaf expansion (From A. Booth, *J. Linnean Soc.* **51**, 166, 1959, and D. Kumar and P. F. Wareing, *New Phytol.* **71**, 639, 1972.)

practices. The earliest commercial use of a plant hormone was the exposure of fruits to ethylene gas to accelerate ripening, a practice which is still common today. Also, we have already mentioned the usefulness of auxins and gibberellins in inducing parthenocarpic fruit development (p. 128). Another practical use for auxins is the induction of flowering, and consequently fruiting, by spraying pineapple crops with compounds such as 2,4-dichlorophenoxyacetic acid (2,4-D). The prevention of "pre-harvest drop" in apples, by synthetic auxin sprays, has also been found to have commercial value. Also, the rooting of stem cuttings is enhanced by treatment with various synthetic auxins (Fig. 5.23).

As we saw earlier (see Fig. 5.5), a characteristic feature of auxins is that above a certain concentration, their effect is to inhibit rather than to stimulate growth. It appears that too high an auxin concentration disorganizes the delicate machinery of growth. Plants treated with an excess of auxin become distorted, with epinastically curled leaves and split stems. They may subsequently die, but not all auxins are equally toxic, and plant species also show varying sensitivities to auxins.

The reason for toxic effects of supra-optimal auxin concentrations is unknown, but an

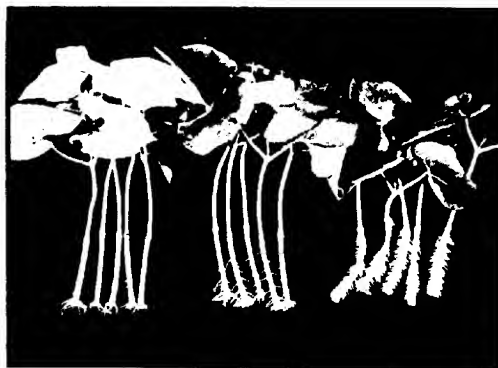


FIG. 5.23. Stimulation of root initiation in bean cuttings. *Left*: control cuttings (no auxin treatment); *middle*: cuttings treated with a solution of 5 mg/l naphthalene acetic acid (NAA); *right*: treated with 50 mg/l NAA. (Original print kindly supplied by Dr. L. C. Luckwill, Long Ashton Research Station.)

important aspect is likely to be the stimulatory effect of high auxin concentrations upon ethylene biosynthesis. Lack of complete understanding of the mechanism of their killing effects has not, however, prevented their use in the control of weeds among crop plants. The removal of weeds by mechanical cultivation is a costly and laborious business, and the discovery that spraying a synthetic auxin, such as 2,4-D, over a field can achieve the same result has proved of enormous economic value. Of course, many chemicals other than auxins, if applied in sufficiently high concentration, are poisonous to plants and can be used as weed-killers. The special merits of synthetic auxins such as 2,4-D are that they will kill plants when applied at relatively low concentrations, they are comparatively harmless to animals, they are non-corrosive, they are translocated within the plant to kill those parts, such as the roots, not reached by sprays and, most important, some auxins are *selective* and can, therefore, be used to kill weeds without damaging surrounding crop plants. As a general rule, cereals are fairly resistant to applied auxins, whereas a number of dicotyledonous weeds are very sensitive and are killed. Thus, the weed known as yellow charlock (*Sinapsis arvensis*) can be controlled in cereal crops such as oat (*Avena*) or wheat (*Triticum*), and lawn-weeds such as daisies (*Bellis perennis*) and plaintains (*Plantago* spp.) can be killed without harming the grass.

The selective action of auxins as weed-killers depends on a number of factors. Very often, susceptible plants are dicotyledonous and possess broad horizontally spreading leaves, which retain auxin solution sprayed on them, while resistant plants are often monocotyledonous with narrow, erect leaves, off which the spray droplets run. In addition, the epidermis of some plants is more easily penetrated by auxin solutions than is that of others.

More important than these factors, though, is the existence of inherent differences between the living cells of different species in sensitivity to synthetic auxins.

Among the most widely used synthetic auxins in weed control at the present time are 2,4-D, 2,4,5-T and MCPA (Fig. 3.8) or proprietary mixtures of these. IAA, the naturally occurring auxin of most plant species, is relatively ineffective as a weed-killer, probably due to its rapid destruction by the enzymes of the treated plants (p. 54).

Gibberellins might have at least as great a practical potential as auxins. The effectiveness of gibberellins in the induction of parthenocarpic fruits is just beginning to be realized at the commercial level, as they are often useful with species in which fruit set is not promoted by auxin. Also, gibberellins act synergistically with auxins in inducing fruit set in species such as tomato. Uses for gibberellins in the cultivation of grapes (*Vitis* spp.) include stimulation of elongation of clusters (to decrease rotting of individual berries), enlargement of berries, and increasing berry set. Strawberry plants can, by treatment with gibberellic acid, be induced to produce earlier flowers, and therefore fruits, than untreated plants. This effect can lead to commercial gain, for fresh strawberries sell for higher prices during the early part of the harvest period. Gibberellic acid has been found to delay ripening of citrus fruits on the tree, and also to improve skin colour in the fruits of certain varieties.

The effectiveness of gibberellins in speeding up germination rates of seeds of a large number of species, as well as in breaking dormancy (p. 270), has obvious practical possibilities.

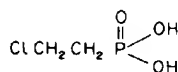
Probably the most important single commercial application of gibberellins yet found is in the malting industry. Malting is a process whereby barley seeds are allowed to germinate for several days. The germinated seeds are then used in the preparation of a medium for fermentation yeasts in the manufacture of beer. The purpose of the malting procedure is to allow time for stored food reserves in the endosperm of each seed to be converted to other substances more suitable as substrates for yeast growth. Thus, for example, starch reserves are converted to sugars by the action of hydrolytic enzymes such as α -amylase which are formed in the aleurone cells during germination (p. 278). Due to the stimulatory effect of gibberellins on α -amylase synthesis (p. 89), treatment of barley seed with gibberellic acid both speeds up and provides more strict control of malting. This has the benefit of conferring considerable saving of time and production of higher yields of the malting product.

A number of plant growth-regulating chemicals in widespread use are grouped together under the general heading of *Growth Retardants*. Thus, commercial preparations known as "B-9" (= "Alar"), "AMO-1618", "Cycocel" (= "CCC" or "Chlormequat"), "Phosfon-D", and "Ancymidol" (= "EL-531") are all growth retardants. Their uses include the inhibition of shoot elongation in ornamental plants (e.g. chrysanthemums, poinsettias, *Coleus*, petunias, etc.) resulting in more compact and desirable plants. Similarly, they are used to shorten and strengthen the stems of some crops, particularly wheat, and thereby reduce "lodging" (the bending over of shoots by heavy rain and wind which can cause severe harvesting difficulties and premature grain germination). There are many other applications of these growth retardants, such as to increase flowering, fruit-set, rooting of cuttings, tuberization, bulbing, resistance to drought, cold and salt. The physiological and

biochemical mechanisms by which these useful effects are achieved are generally ill-understood, but at least some of them result from the capacity of growth retardants to inhibit gibberellin biosynthesis in plants (see p. 58). There are a few other non-phytotoxic growth inhibitory chemicals used in agriculture and horticulture, such as maleic hydrazide, but these do not appear to act by affecting gibberellin biosynthesis.

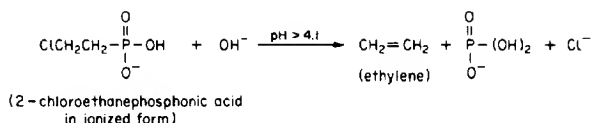
It is highly likely that some synthetic cytokinins will be widely used in future to delay the processes of senescence in flowers and vegetables whilst these products are being transported from the grower to the consumer.

As already stated, ethylene has been used for some years for accelerating ripening processes in a number of fruits (an aspect of senescence), particularly citrus fruits such as oranges and lemons. Moreover, demonstrations that this gas is a most potent stimulator of premature leaf fall have raised the possibility of its use as a defoliant in crops such as cotton, pea, beans, etc., where mechanical harvesting of fruits is expedited if leaves are not present. However, because of its gaseous nature, ethylene does not readily lend itself to applications to outdoor crops. Because of this, a search has been made for other chemicals which promote leaf abscission. A number of such substances are now known, and the most effective of these are those which enhance ethylene synthesis in the plants to which they are applied. A recent development is the search for chemicals which can be sprayed on to plants, but which are themselves broken down in plant tissue to yield ethylene. An example of such a substance is 2-chloroethanephosphonic acid, known commercially as "Ethrel" or "Ethepon":



2-chloroethanephosphonic acid

This compound undergoes chemical decomposition at pHs higher than 4.1. Because plant cells normally have a pH above 4.1, an aqueous solution of Ethrel will on entering the tissues be degraded, liberating ethylene:



Ethrel therefore induces physiological responses in plants identical to those elicited by ethylene, and is gradually being introduced to agriculture and horticulture. At the present time it is already in widespread use to increase the rate of latex flow from rubber trees, and

to hasten fruit ripening and leaf abscission. It is also coming into use to control branching and suckering of both crop and ornamental plants.

There is little doubt that practical uses of plant growth hormones will in the future be more extensive and varied than at present. Full realization of their potential will, however, not be achieved until our understanding of their functions and modes of action in plants is greatly increased.

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CHAPTER 6

Aseptic Culture Methods in Studies of Differentiation

HISTORICAL ASPECTS OF PLANT TISSUE CULTURE

In the preceding and later chapters of this book various aspects of morphogenesis are considered and a number of experimental approaches to the subject are described. No one aspect of botanical research is able to provide a unified picture of differentiation, and studies of this subject must aim at the unification of information derived from the fields of morphology, physiology, biochemistry and biophysics. This is a difficult task, for in the intact plant there are complex interactions between the various processes underlying growth and differentiation. Consequently, it is desirable to reduce the complexity of the system so that the controlling processes may be more easily identified and studied. This can be done to varying degrees, by isolating embryos and other parts of a plant in order to eliminate certain complicating correlative influences during studies of their behaviour. In general, isolated embryos and plant parts must be carefully nurtured to keep them alive and free from infection by micro-organisms. This normally entails the use of *aseptic culture techniques*. Such methods often facilitate the maintenance of isolated embryos, organs and tissues for considerable periods of time. However, the culture of isolated plant tissues, organs and embryos presented many difficulties to earlier workers which were only gradually surmounted.

The smallest viable unit of a plant one can at present envisage as reproducing, growing and developing in culture is a single cell. As long ago as 1902, Haberlandt attempted to grow single plant cells in aseptic culture, but, for various reasons which we now understand, his attempts were unsuccessful.

Following Haberlandt, other workers established methods which would allow the growth of isolated plant organs and tissues in culture. Excised roots were the first plant organs to be successfully brought into aseptic culture, and work by White in the 1930s demonstrated that given appropriate nutrients, such roots would grow and differentiate normal root tissues. Work by Gautheret (1939) and others established that isolated portions of storage tissue, e.g., from carrot roots, could be kept alive and grown in aseptic culture. *Callus cultures* derived from such isolated tissues lend themselves to studies of the effects of nutrients, vitamins and hormones upon cell division, differentiation of vascular tissues, and

the inception of organized meristematic regions within the predominantly parenchymatous tissue. By definition, a callus is a mass of proliferating tissue consisting predominantly of parenchymatous cells, but in which differentiation may occur under suitable conditions.

When a callus is grown in agitated liquid culture, cells at the surface are often broken away and float free in the medium to give a *liquid suspension culture*. Usually, such free cells do not divide when retained in a medium suitable for callus growth, for propagation of free plant cells demands a more elaborate medium than does a callus. Media have, however, been devised which are capable of supporting proliferation of free plant cells.

ORGAN CULTURE

Root Culture

It has proved possible to grow several types of plant organ in aseptic culture, including roots, shoot apices, leaves, flower parts and fruits. The nutrient requirements for such organ culture vary considerably from species to species and according to the type of organ in question, but certain general requirements can be recognized. Intact higher plants are autotrophic; that is, they are able to synthesize all the organic substances required for their own life from carbon dioxide, oxygen and mineral nutrients. However, since most aseptic cultures are unable to carry on photosynthesis, it is clear that they will require at least a carbon source, usually supplied in the form of a sugar such as sucrose or glucose. In addition, aseptic cultures require the same mineral nutrients as the intact plant, including both macronutrients (nitrogen, phosphorus, potassium and calcium) and micronutrients (Mg, Fe, Mn, Zn, etc.).

In addition to those requirements for a carbon source and mineral nutrients, it is found that most isolated organs have also a requirement for certain special organic substances. Thus, most isolated roots grown in aseptic culture require to be supplied with certain vitamins, particularly thiamin (vitamin B₁) and sometimes pyridoxin (vitamin B₆), nicotinic acid and others. It appears that in the intact plant, certain vitamins are synthesized in the leaves and that roots are dependent upon the shoots for the supply of these substances, which they are unable to make for themselves. Tomato roots require only sucrose, mineral nutrients and thiamin, and given these they will grow successfully in culture for many years.

Excised roots of some monocotyledonous plants fail to grow even when supplied with a full complement of B-vitamins and other vitamins. In some of these cases (e.g., rye), an exogenous auxin supplement to the nutrient medium allows growth to proceed. In order to maintain the culture, the excised roots must be regularly subcultured on to fresh medium, by excising a piece of root bearing a lateral, which then proceeds to grow rapidly and maintain the culture.

Excised roots of most species produce only root tissues in culture, but there are some species the cultured roots of which regenerate shoot buds as well as further roots, e.g., *Convolvulus*, dandelion (*Taraxacum officinalis*) and dock (*Rumex crispus*).

Culture of Shoot Apices and Leaves

Isolated shoot apical meristems and leaf primordia can also be grown in aseptic culture. These frequently produce adventitious roots and can eventually develop into complete plants. The shoot apices and leaves of vascular cryptogams, such as ferns, are relatively more autotrophic than those of angiosperms. Thus, even a small fern apex can be grown on a medium containing only a carbohydrate source and mineral nutrients. Small angiosperm apices (less than 0.5 mm in diameter) require a general source of organic nitrogen and certain specific amino acids and vitamins, in addition to the basic medium, but larger apices will grow on a simple medium. The simpler requirements of large apices may be due to the fact that they carry larger leaf primordia, which apparently can supply some of the requirements of the apex for vitamins and other organic nutrients.

Isolated young leaves of the fern, *Osmunda cinnamomea*, and of sunflower (*Helianthus annuus*) and tobacco (*Nicotiana tabacum*), have been successfully grown on a simple medium containing only sucrose and inorganic salts (Fig. 2.12). Such isolated leaf primordia continue to grow and develop into normally differentiated leaves, although they are usually very much smaller than normal leaves developed on the plant (p. 36).

EMBRYO CULTURE

We have earlier (p. 24) given an account of the development of embryos as it occurs naturally within the embryo sac. Experimental approaches to the study of plant embryogenesis have in recent years made extensive use of aseptic culture of isolated young embryos.

The easiest method of culturing an embryo is to allow it to develop *in situ* within the ovary or dissected out ovule. The ovule, if placed on a suitable nutrient medium, is able to support the development of the zygote to maturity. The presence of placental tissue aids the development of the ovule and embryo, and consequently it is easier to maintain ovules in aseptic culture when they are left within the ovary. The physiological requirements of ovules do not appear to be species specific, since young fertilized ovules of widely different species have grown to mature seeds following transplantation on the placenta of *Capitum* fruits.

Although embryos can be grown from the zygote stage to maturity when they are left inside a cultured ovule, they show complex nutrient requirements when isolated from the ovule. The mature embryo is, of course, autotrophic and will grow if provided with the normal conditions necessary for germination, viz. adequate water and oxygen and a favourable temperature. However, if embryos are excised at younger stages it is found that they will not grow, even if supplied with sucrose and mineral salts.

A useful technique for the culture of young embryos was introduced in 1941 by van Overbeek, who found that they could be grown from a quite immature stage by supplementing sucrose and mineral salts with coconut milk, which is a liquid endosperm.

This endosperm allows the development of the coconut palm embryo, as it contains a complex range of substrates necessary for the growth of the embryo, and it is also very effective in supporting the growth of embryos of other species. Attempts have been made to identify the active components of coconut milk and they are now known to include sugar-alcohols such as myoinositol, leucoanthocyanins, cytokinins, and probably also auxins and gibberellins. By using this technique it has been possible to develop to maturity embryos of certain species isolated at an early stage of development.

As embryos develop they appear to become less heterotrophic, as has been demonstrated by observations on isolated embryos of *Capsella* and *Datura*, which showed that globular embryos did not survive at all in culture, but that early heart-shaped embryos (see Fig. 2.2) would develop further if supplied with a nutrient medium containing sugar, inorganic salts, vitamins and coconut milk. With slightly older heart-shaped embryos the coconut milk may be replaced by a source of reduced nitrogen such as L-glutamine, and with more highly developed embryos even the glutamine can be omitted. Thus, there appears to be a progressive increase in the synthetic abilities of the embryo during its development.

On the other hand, there is more recent evidence that the concept of a decrease in heterotrophic nutrition with increase in embryo age may not be strictly true. Even young globular embryos of *Capsella* have been cultured to maturity in a relatively simple medium containing no organic nitrogen source such as L-glutamine, nor high sucrose concentration. But, in addition to the usual inorganic salts, vitamins and 2 per cent sucrose, the medium did have to contain a balanced mixture of very low concentrations (about 10^{-7} M) of an auxin (e.g., IAA), a cytokinin (e.g., kinetin) and adenine sulphate. It appears that, in the case of *Capsella* at least, it is the balance of growth hormones in the immediate environment rather than substrate level metabolites, which plays a determining role in embryogenesis.

TISSUE CULTURE

In contrast to the techniques of organ culture, tissue culture involves the aseptic culture of an isolated homogeneous mass of cells. All plant organs consist at the time of their excision of a number of different tissue types, and therefore represent more complex systems than do isolated individual tissues. In studies of the physiology and biochemistry of morphogenesis it is desirable to work with as simple a system as possible, and clearly culture of isolated tissues would appear to represent a simplification of experimental material in comparison with organs in culture. Fortunately, tissues from many sources can be maintained in culture for an indefinite period, and afford enormous but as yet largely untapped, possibilities for research in physiology and biochemistry. Such cultures have already proved very valuable for certain biochemical studies. For example, cultures of sycamore (*Acer pseudoplatanus*) cambial tissue have been extensively used for studies on cell-wall metabolism, and we shall see below several examples of the value of tissue culture for studies on differentiation.

When small pieces of root phloem parenchyma of wild carrot (*Daucus carota*), or pith parenchyma of tobacco (*Nicotiana tabacum*) stem, or even chlorophyll-containing palisade

cells from leaves of *Arachis hypogea* and *Crepis capillaris* are placed on a suitable medium, they can not only be kept alive but can be induced to grow. That is, mature parenchymatous or mesophyll cells, which, if left undisturbed in the plant body, would undergo no further cell division, can be made to divide mitotically, giving rise to an undifferentiated callus. An extreme example of retention of the capacity for cell division in mature plant cells was provided by cultivation of a callus from medullary ray tissue excised from a region adjacent to the pith in 50-year-old lime (*Tilia*) trees. These cells had matured a full half-century earlier, and yet their continued potential for active cell division was revealed under suitable conditions in culture.

It seems likely, therefore, that any living, nucleated, plant tissue can give rise to a proliferating undifferentiated callus when excised and placed on a suitable culture medium. However, great difficulty is often experienced in establishing a callus from a previously untried source of tissue, for there is apparently considerable diversity in nutritional requirements of tissues from different species, or even from different locations within one plant. In general, it has proved easier to culture tissues consisting originally of non-green parenchyma, such as phloem or pith parenchyma. The establishing of green, photosynthesizing, callus growths from chloroplast-containing leaf cells has come much later.

In addition to the usual macro- and micro-inorganic nutrients, and an organic carbon source, isolated tissues are frequently found to require (1) an organic source of reduced nitrogen, which may be supplied as amino acids or, in some species, as the amide of glutamic acid, L-glutamine; (2) vitamins, including thiamin, nicotinic acid and pyridoxine, and (3) the sugar alcohol, myo-inositol. In addition, an auxin, such as 2,4-D, and sometimes a cytokinin, are required. The fact that it is necessary to supply hormones to callus cultures, whereas they are not normally required by organ cultures, may indicate that the organized meristems of organ cultures may be centres of hormone biosynthesis, whereas the parenchymatous tissues from which callus cultures are derived do not have the capacity for hormone synthesis.

It is of interest to note that photosynthetic callus growth from palisade mesophyll cells of *Arachis hypogea* does not need an external supply of any vitamins. This can be related to what was said earlier concerning the normal production of vitamins in the leaves of plants, and their supply to other regions such as the roots.

Repeated sub-culturing of some tissue cultures, such as those of carrot, grape, *Scorzonera*, tobacco, and other plants, leads to a spontaneous and irreversible change in that they acquire the capacity to synthesize excess quantities of auxin. Thus, a tissue which when first brought into culture requires an exogenous supply of auxin and cytokinin, later on, after sub-culture, becomes autotrophic for these hormones. Such long-established callus cultures are then said to be *habituated*, or *energized*, and closely resemble tumorous as opposed to normal plant tissues. For example, plants infected with the crown-gall bacterium (*Agrobacterium tumefaciens*, synonym *Phytophthora tumefaciens*) exhibit tumorous (callus-like) growths at the points of infection. By appropriate heat treatment, the bacteria can be killed and removal of a portion of one of these treated tumours into aseptic culture leads to the production of a massive undifferentiated callus, which is completely self-sufficient in auxin

and cytokinin. Thus, infected or habituated cells have undergone a permanent change in that they are able to synthesize substances which they were unable to produce before. This capacity is transferred from one cell generation to the next and is brought about by transfer of bacterial DNA to the plant cell (see p. 312).

SUSPENSION CULTURE OF PLANT CELLS

A suspension culture consists of cells and small aggregates of cells dispersed and growing in a moving liquid medium. The principal problem encountered initially in attempts to make isolated single cells and small aggregates of cells divide in culture is that such isolated cells are "leaky". For a number of reasons, particularly their large surface areas exposed to the liquid medium, in contrast to callus cells surrounded by like cells, isolated plant cells in an agitated liquid medium tend to lose substances required for cell division to the medium. Consequently, the nutritional requirements of free plant cells and small cell aggregates may be more complex than those for callus cultures of the same species, since it is necessary to supply the substances which tend to be lost by the cells into the medium. For some types of cell culture it has been possible to determine the precise nutrient requirements so that they can be grown on a defined medium, which usually included the various constituents already listed for callus cultures (p. 145). In other cases, it has not yet been possible to grow cell culture in a defined medium and it is necessary to add coconut milk, which must, therefore, include certain, as yet unknown, special nutrient factors.

Although all the nutritional problems involved in growing colonies of cells from different sources have not yet been solved, we are able now to see the general picture. What is striking is the very close similarity between nutritional requirements for suspensions of plant cells and for development in isolated young embryos (p. 144). In the intact plant these requirements are met by surrounding tissues—particularly, in the case of embryos, by the endosperm.

Plant cells growing in suspension culture look very much alike from whatever species they originated. The principal characteristics of cells in suspension culture are: (i) numerous and large vacuoles, even in cells capable of division; (ii) prominent cytoplasmic strands which show active streaming movements; (iii) a large nucleus with nucleolus (Fig. 6.1 *Left*). In addition, a variety of cell types coexist in a given cell suspension, only some of which are free single cells. Some free cells divide and give rise to clusters of smaller, more dense cells. Others increase in size and divide with the formation of internal cross-walls to produce either a filament of cells or, in some cases, a new free cell by a process analogous to "budding" of yeast cells in culture (Fig. 6.1 *Right*). Thus, cells of higher plants in suspension culture do not have the morphology of cells in the tissue from which they were derived. Furthermore, they evidently have a different pattern of metabolism, for they usually do not contain typical storage products.

Ideally, the initial inoculum of isolated cells in a culture vessel will consist exclusively of individual free cells. Only rarely is this ideal attained, by, for example, filtering the suspension through a sterile gauze possessing a pore diameter too small to allow the passage of cell

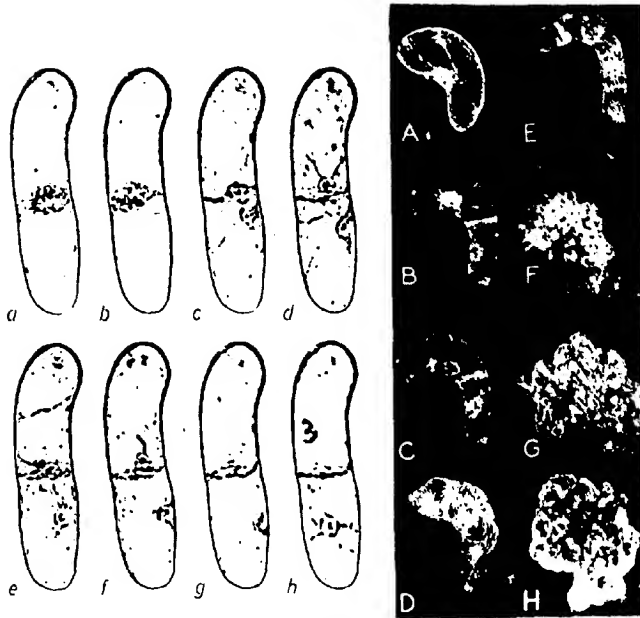


FIG. 6.1. *Left:* Cell division in a single isolated cell from *Phascolus vulgaris*. Note the large vacuole, prominent cytoplasmic strands and large nucleus with nucleolus. Time lapse between pictures a, b and c, 30 minutes; between c, d, e, f, g and h, 60 minutes. (Reprinted by permission of the Rockefeller University Press from the *Journal of General Physiology*, 43, 843, 1959-60. Print supplied by Dr. Ludwig Bergmann.)

Right: Development of a cluster of cells from a single cell of tobacco stem-pith isolated in aseptic culture. A. Single cell 1 day after placing in culture medium. B-H. Stages in the formation of a mass of cells, from the single cell in A. (From W. Vasil and A. C. Hildebrandt, *Science*, 150, 889-92, 1965. Print supplied by Dr. A. C. Hildebrandt.)

clusters. By such means a very dilute suspension of free cells (5 or less cells per cm^3 of nutrient medium) can be set up. Under suitable conditions this suspension of cells will multiply so that in 2 or 3 weeks' time the cell density will have risen to approximately 100,000 per cm^3 . Microscopic examination of the cell population at this time reveals that not all the cells are now free—i.e. cell clusters of various sizes and shapes are usually present in addition to single cells. Various studies have shown that formation of a multicellular cluster in a suspension culture of plant cells takes place by repeated division of one cell, the daughter cells of which do not separate. Separate free plant cells do not aggregate into clusters in the way that some cultured animal cells do. In most research with cell suspension cultures, the original inoculum contains not only free cells but also small aggregates of cells and even dead cells and cell debris.

REGENERATION STUDIES WITH ASEPTIC CULTURE

Plants show remarkable capacities for regenerating whole organisms from isolated pieces of shoot, root, or leaf, and even from relatively unorganized tissue such as callus. Some more general aspects of regeneration will be discussed later, but we shall first describe experiments on regeneration which have been carried out with callus and suspension cultures.

In callus cultures cell division occurs randomly in all directions and gives rise to an unorganized mass of tissue; thus, there are no clearly defined axes of polarity in a callus. By contrast, in a shoot or root meristem we have a highly organized tissue structure, in which, as we have seen, quite well-defined patterns of division can be recognized. It has been found that under certain cultural conditions shoot and root meristems can be formed within a callus, so that whole new plants may be regenerated.

Root and Bud Regeneration in Callus Cultures

We have already made mention in Chapter 3 (p. 60) of Skoog's studies on the interacting influences of auxins and cytokinins on the growth of tobacco pith-derived tissue cultures. Skoog observed that cytokinin and auxin interacted to initiate cell division, but he also found that these same two types of growth hormone could interact to initiate organized meristems. Thus, it was discovered that if the *proportions* of auxin and cytokinin were varied, then the pattern of meristem formation was altered. When the proportion of auxin to cytokinin was relatively high, there was differentiation of some callus cells into root primordia. A higher concentration of cytokinin relative to auxin caused cells to differentiate into shoot apical meristems. Subsequent growth of the root and shoot primordia led to the callus cultures having the appearance shown in Fig. 6.2. Thus, small changes in the auxin-cytokinin ratio could (a) initiate meristems and (b) channel differentiation of these into either shoot or root apical meristems.

Control of root or shoot-bud formation in callus cultures by variations in auxin-cytokinin balance has now been demonstrated for tissues of several origins. The interacting effects of the hormones in this phenomenon can be modified by other factors, such as sugar and phosphate levels, sources of nitrogen, and other constituents of the medium such as purines. There is, however, no doubt that auxin and cytokinin may regulate not only the initiation of organized meristematic centres in the callus, but also the type of meristem formed. Nevertheless, stimuli other than auxin and cytokinin are involved in apical meristem initiation in callus cultures. For example, initiation of lateral roots in pea-stem segments is inhibited by red light. Thus a phytochrome-based mechanism (p. 183) may be involved in the initiation of root apical meristems.

Embryo Formation in Aseptic Cultures of Plant Cells and Tissues

Totipotency of plant cells has perhaps been most spectacularly demonstrated by the regeneration of whole plants from embryos formed in cultures of both somatic cells and

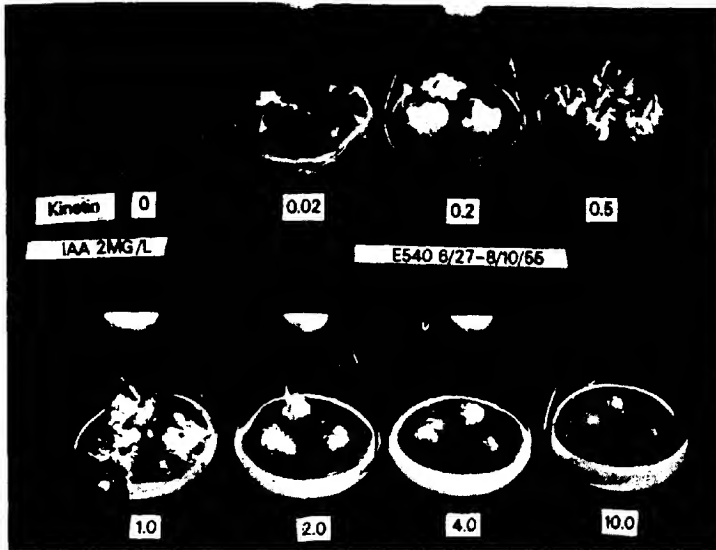


FIG. 6.2. Effect of a range of concentrations (0-10 mg/l) of kinetin on growth and organ formation in tobacco-pith derived callus cultured on nutrient agar containing, in all flasks, 2 mg/l indole-3-acetic acid. All cultures 44 days old. (Reprinted from *Symp. Soc. Exp. Biol.* **11**, 1957. Print kindly supplied by Professor F. Skoog.)

those of male generative cells (i.e. pollen grains). The formation in tissue cultures of plant embryos of at least superficially normal appearance and behaviour was first observed by Steward in 1958 and Reinert in 1959, who by imposing a sequence of changes in the composition of the nutrient media caused callus cultures of carrot-root phloem parenchyma to give rise to embryos which were very similar to normal embryos and which on transfer to a suitable medium developed into whole carrot plants. Embryos formed from cells of origins other than a fertilized egg are often referred to as either *adventive embryos* or *embryoids*. The changes in the nutrient media required to bring about adventive embryo formation principally involved alterations in the balance of auxin and cytokinin.

Since that first report, other workers have obtained adventive embryos in callus cultures, in suspension cultures, and in cultures of isolated anthers and pollen grains (Fig. 6.3). Regeneration in cultivated and wild varieties of carrot (*Daucus carota*) has been extensively investigated, particularly with respect to embryogenesis in aseptic culture (Fig. 6.5), but it has been found that the capacity for adventive embryo formation is widely distributed in the plant kingdom. Nevertheless, one cannot assume that all cells of a plant, or all species, retain their totipotency, for numerous unsuccessful attempts have been made to obtain embryogenesis, and/or organogenesis, in aseptic cultures of tissues from many species. It is

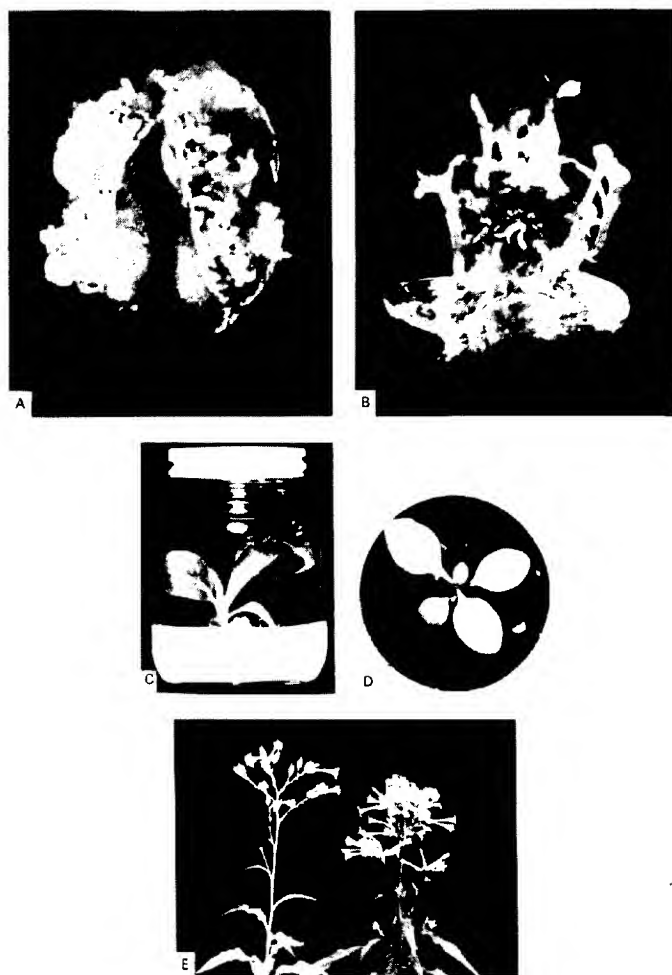


FIG. 6.3. Development of embryoids (adventive embryos) from pollen grains during anther culture of *Nicotiana tabacum* c.v. "White Burley". A. Anther cultured for 28 days at 25°C in continuous light, showing a large number of germinating embryoids, each of which has developed from one pollen grain. B. The same anther 7 days later. At this stage the plantlets can be readily teased apart and transplanted individually as shown in C. The transplanted plantlets can later be transferred to a mixture of peat and sand in a normal plant pot (D). E. Inflorescences of, on the left, a normal diploid tobacco plant, and on the right, of a haploid plant grown from a pollen grain as shown in A–D. It is interesting to note that haploid plants usually develop larger inflorescences than do diploid plants. (From N. Sunderland, "Pollen and Anther Culture", Chapter 9 in *Plant Tissue and Cell Culture*, ed. H. E. Street, Blackwell Scientific, 1973. Original prints supplied by Dr. N. Sunderland.)

possible that only true diploid cells (or haploid in the case of pollen grains) can undergo embryogenesis, and that failures to induce adventive embryo formation may have been associated with polyploidy in cultured cells and tissues.

The formation of an adventive embryo is marked by the appearance of an organized group of cells possessing longitudinal polarity and also with, at a very early developmental stage, a shoot and a radicular pole at opposite ends. The adventive embryo is formed without any connection with vascular tissues of the mother plant or callus, and this contrasts with the pattern of formation of monopolar buds and roots, as these are always connected to such vascular tissue.

Adventive embryo formation can, under appropriate experimental conditions, be a continuous process, and a culture may therefore contain, side by side, embryos of all stages of development. Adventive embryos induced in tissue cultures appear to be formed from single cells, usually at the upper surface of the callus. The surrounding callus cells may act as a "nurse-tissue" to the embryo in its very early stages of development. There is some evidence that the formation of adventive embryos in suspension cultures of plant cells similarly involves a nurse-tissue, in that there is first the production of a multicellular aggregate (i.e. a small callus) by repeated division of an originally single cell, following which embryos may be initiated from single cells at the surface of the aggregate. On the other hand, it has been claimed by Steward and his co-workers that isolated free single cells can develop directly by segmentation without callus formation to pro-embryos and that these then go on to form embryos and normal plantlets. Steward has, in fact, argued that one of the requirements for the initiation of adventive embryos in suspension cultures is that cells must be set free from associations with their neighbours and thus be able to grow independently. However, various studies in recent years have shown that adventive embryos can form from individual cells at the surface of a callus. More microscopic studies of adventive embryos will undoubtedly resolve the question as to whether or not a single plant cell, not in contact with other cells, can develop directly into an embryo. Recent electron-microscope studies have, however, indicated that embryogenic cells in callus from *Ranunculus scleratus* possess protoplasmic strands, linking them with adjacent callus cells during embryogenesis. This does indicate that cells do not have to be physiologically isolated from other cells for their development into an adventive embryo, and in addition strengthens the suggestion that nurse-cells are in some way necessary in early embryogenesis. Similarly, the concept of nurse-cells is supported by the reports that adventive embryos form in suspension culture from one cell of a cell aggregate, with the remaining cells of the aggregate in some way supporting the initial development of the embryoid. Nevertheless, it is true that only "exposed" cells of a callus, or a cell in suspension culture, actually become embryogenic, even though they may require the support of "nurse-cells", and that the mass of cells in a callus do not form embryoids but may do so if they are separated into a suspension culture.

Many of the earlier experiments on adventive embryo formation in aseptic culture involved the use of media containing the liquid endosperm of coconuts. At one time it was considered that coconut milk contained special substances which were essential for the

initiation of embryos in suspension cultures. More recent research has, however, shown that cells in suspension culture can form embryos on synthetic, chemically defined, media in the absence of liquid endosperm. Thus, one does not need to invoke unknown nutritive or hormonal "embryo-inducing" factors in liquid endosperms. Adventive embryo formation is achieved experimentally by successive changes in the nutrient media, an important aspect of which includes the nitrogen to auxin ratio. Because of this a number of workers utilize a sequence of media changes which culminates in transfer to an auxin-free from an auxin-containing medium, and this can be sufficient to induce embryogenesis in the culture.

Pollen- and Anther-Culture

Over the past few years, the techniques previously developed for adventive embryo formation from somatic cells have been used to produce embryos and even entire whole plants from pollen grains. Pollen grains are produced naturally in relatively large numbers and in an easily accessible form for the research worker. Each grain consists of just a few cells (five in gymnosperms; three in angiosperms) and possesses a unique genome derived from the process of meiosis in microsporogenesis. In true diploid species each gene in the pollen grain is present as a single copy only, and therefore formation of an adventive embryo from the pollen grain results in a haploid plant in which every gene can be expressed in the phenotype. This fact has extremely important connotations in the practice of plant breeding, for experimentally produced mutant genes can be evaluated very much more quickly and easily in haploids. In addition, the exposed adventive embryos from pollen or anther cultures and derived plantlets also lend themselves to convenient and uniform mutation-inducing treatments such as irradiation with X-rays.

The techniques required to induce embryogenesis in pollen and anther cultures are still in the process of development, and to date only a limited number of species have been successfully propagated in this way. The first experiments on pollen culture took place in the 1950s, when Tulecke was able to cause pollen of certain gymnosperms to proliferate into callus. Not until the mid-1960s was angiosperm pollen brought successfully into aseptic culture by Guha and Maheshwari, and then only by culturing the whole anther of *Datura innoxia*. However, Guha and Maheshwari also found in 1967 that development in cultured anthers of *D. innoxia* led to the formation of haploid plants, with one pollen grain giving rise to one plant. Since that time aseptic cultural conditions have been developed which permit growth and embryogenesis in pollen (usually within the excised anther) in various other angiosperms (Fig. 6.3).

Methods of anther culture usually involve aseptic excision of anthers, and their transfer to the surface of sterilized agar medium, or flotation on the surface of liquid medium, or onto filter-paper bridges over liquid medium. An important aspect of the procedure is the selection of anthers containing pollen at an appropriate stage in microsporogenesis. Maximum yield of adventive embryos in *Nicotiana tabacum* anther culture is obtained when

the excised anthers contain pollen at the stage when the vegetative cell is in the process of rapid cytoplasmic synthesis, but results obtained with other species have indicated that the critical stage may vary from one species or cultivar to another. Furthermore, it may also be influenced by aseptic culture conditions and the environment under which the donor plants were grown. Culture media used in pollen and anther culture are similar to those used in the aseptic culture of somatic tissues, and as with these it is the hormonal component of the medium which is critical. For anthers of most species, both auxin and cytokinin are included, although in a few instances either one or the other is sufficient. Cytokinin can be replaced, if desired, by coconut milk.

Embryogenesis in angiosperm anther culture occurs by development of the vegetative cell of each pollen grain. The initial stages of adventive embryo formation varies from species to species. In some cases the vegetative cell divides repeatedly, and there is complete suppression of pollen-tube development. Gradually the derivatives of the vegetative cell can be seen to be organized as an embryo which further develops into a plantlet. In other cases, embryogenesis appears to occur where the normal unequal division resulting in a vegetative and generative cell is replaced by an equal division (p. 319).

Under certain cultural conditions, and depending on species, pollen in cultured anthers may also divide in manners which give rise to a callus. Shoot buds and root initials often form from such a callus and further plantlets can be obtained in this way. Although much research remains to be done, it is already clear that pollen and anther culture techniques will be very important in future breeding of new crop and ornamental plants.

Isolation and Culture of Plant Protoplasts

Since the mid-1950s methods have been devised which allow the isolation of protoplasts from somatic and reproductive cells. A protoplast is, of course, a plant cell from which the cell wall has been removed, and in aseptic culture isolated naked protoplasts behave in many ways rather like animal cells in culture. Under suitable cultural conditions they grow and divide, but can also be induced to regenerate cell walls.

Plant protoplasts can be isolated from their parent tissues by either mechanical or enzymatic means. Mechanical isolation of protoplasts has been achieved by first plasmolysing the cells in a hypertonic plasmolyticum and then carefully cutting away the walls by microsurgical procedures. Such a technique therefore cannot be used on non-vacuolated cells such as those of meristems, and for this and other reasons enzymatic methods of protoplast isolation have been adopted by many workers in this field. The enzymes used are ones which degrade components of plant cell walls, and most commonly a pectinase (to separate cells) and a cellulase (to degrade the cellulosic walls) are used either sequentially or as a mixture. Once isolated, plant protoplasts have to be kept in a liquid culture medium, the osmotic potential of which closely matches that of the protoplasts; otherwise irreversible damage can be caused by the protoplasts bursting or shrinking excessively.

Isolated protoplasts in aseptic culture have very great potential uses to research workers.

For example, uptake of viruses and various macromolecules by plant cells is very much more readily studied in the absence of cell walls. However, a most exciting prospect lies in the successful fusion of protoplasts of differing origins to produce new plant species by somatic hybridization. The breeding of new plants has always been restricted by the necessity of effecting sexual fertilization. Failure to obtain viable hybrid embryos from interspecific and intergeneric crosses can be due to a number of causes, arising from disruption of the normal sexual processes, e.g., by failure of the pollen tube to penetrate the embryo sac, breakdown of the endosperm, etc. Such difficulties could be avoided if sexual reproduction can be by-passed by direct fusion of vegetative cells. Successful fusion of isolated plant protoplasts has already been achieved (Fig. 6.4), and appropriate methods have already been devised to allow some fused protoplasts to regenerate a whole new plant, either by adventive embryo formation or by root and shoot-bud initiation from a derived callus.

In addition to providing a means of circumventing natural incompatibility mechanisms in genetic hybridization, the mixing of the cytoplasm and organelles of different plant species is also of potential, but as yet unrealized, practical importance. For example, one possible development would be the fusion of protoplasts between different divisions of the plant kingdom, which may, for example, provide one means for the creation of cereal plants containing a characteristic of blue-green algae in being able to utilize atmospheric nitrogen rather than being dependent on nitrogenous fertilizers added to the soil. The fusion of plant and animal cells has been already achieved, and there is the possibility that such procedures could lead to the creation of plant-animal hybrids.

REGENERATION IN SHOOT AND ROOT CUTTINGS

Although not normally performed in aseptic culture, one of the most obvious examples of regeneration is seen in the common horticultural practice of vegetative propagation of plants by taking shoot or root cuttings, and allowing them to develop adventitious roots and/or buds. In shoot cuttings, a callus is frequently formed at the base of the cutting, as a result of divisions originating in the cambium, and from such a callus root primordia arise. However, adventitious roots may also be formed in normal tissues of the stem—usually in the pericycle, but in some species in the cambial zone. In root cuttings, both roots and buds commonly arise from callus formed from parenchyma in the younger phloem.

The ease with which roots can be formed on shoots varies enormously: cuttings from plants such as bean will produce roots if simply left with their lower ends immersed in water, whereas those from other species will do so only rarely, even under what appear to be the most favourable conditions. In those species which do produce roots, it is generally true that a piece of stem which possesses a bud or leaves will form roots at its base, but that a disbudded and defoliated stem piece produces roots much less readily or not at all. This suggests that a substance is formed in the buds and leaves which moves downward and stimulates root formation at the base of the stem. The existence of such a root-initiating substance in young leaves has been proved by the demonstration that extracts of young

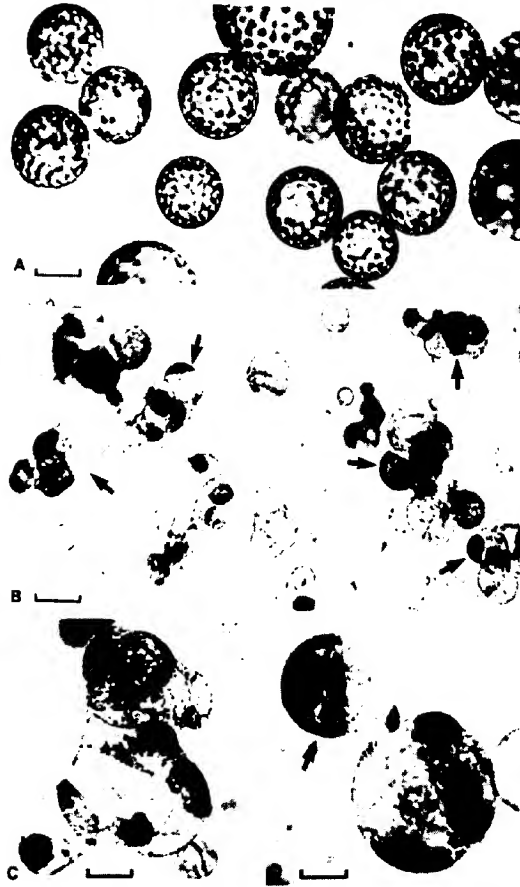


FIG. 6.4. Isolated plant cell protoplasts and inter-species fusion. A. Protoplasts isolated from leaf mesophyll of *Petunia hybrida* following treatment of the leaf with cell-wall-degrading enzyme (bar represents 25 μm). B. Protoplasts isolated from leaf mesophyll of *Petunia hybrida* and colourless leaf epidermis of *Nicotiana tabacum* which have been treated with polyethylene glycol to induce aggregation and fusion. Several fusing inter-species aggregates are visible (arrows) (bar represents 50 μm). C. Large aggregate containing fusing protoplasts of *Petunia hybrida* and *Nicotiana tabacum* (colourless epidermal protoplasts) (bar represents 25 μm). D. Inter-species heterokaryon which has resulted from the fusion of several protoplasts of *Petunia hybrida* and of *Nicotiana tabacum*. The heterokaryon has rounded off and extensive mixing together of the cytoplasm is beginning to take place. Two very closely adhering protoplasts of the two species are also visible (arrow) (bar represents 20 μm). (Original prints supplied by Professor E. C. Cocking.)

leaves do stimulate the rooting response of stem cuttings. Further, it has been found that the application of auxins to cuttings has a similar effect to that of extracts of young leaves (Fig. 5.23), suggesting that the stimulatory effect of buds and leaves upon rooting is probably due to the production of auxin by these organs. The effect of auxin in rooting of cuttings is to increase the rate of formation and absolute number of adventitious root initials. This is, therefore, another example of cell division and differentiation being activated by auxin.

The formation of roots in stem cuttings normally occurs at the *basal end* of the stem. This is true even if the cutting is inverted, so that the morphological lower end is uppermost (p. 331). The stem therefore shows polarity in the initiation of roots. The fact that auxins are known to stimulate the formation of root initials and also that auxins move in a basipetal manner (p. 99), lead one to believe that the polarity shown in root formation is a consequence of the movement of auxin to the morphologically lower tissues, where its arrival triggers off the processes of root initiation. In fact, if an auxin is applied to the apical end of a stem cutting, then callus formation and subsequent root formation is stimulated at the base of the cutting. If applied basally, then roots are again stimulated there.

If a stem section is dipped in a solution of cytokinin it may react by producing many buds at the morphologically upper end of the stem, but few or no roots, the opposite effect to that elicited by dipping in an auxin solution. Nevertheless, as we have seen (p. 148), the stimulatory effect of auxins upon root formation may not be revealed unless the responding tissues also contained an appropriate concentration of cytokinins, since root formation involves active cell division. Root cuttings behave similarly to stem cuttings with regard to polarity of root and shoot bud initiation, and the effects of auxins and cytokinins (p. 331). The fact that buds and roots are formed at opposite ends of isolated segments of stem or root appears to be the result of movement of auxin and cytokinin in opposite directions, a preponderance of one or the other accumulating at either end, causing either buds or roots to be initiated. Indeed, it has been found that if cuttings of chicory (*Cichorium intybus*) roots are placed under moist conditions, which favour regeneration, then certain changes occur in the distribution of the endogenous hormones within the cuttings, so that high auxin concentrations are found at the basal ends, and high cytokinin levels at the apical ends. These changes occur *before* there is any observable regeneration of buds and roots and hence they may play an important role in the pattern of regeneration. Certainly these observations are consistent with the findings that bud and root regeneration in callus cultures are associated with high cytokinin and high auxin levels, respectively (p. 148).

At the present time we know little of the translocation patterns of cytokinins in plants. Kinetin itself is apparently not translocated readily, for it remains at, or very close to, the place to which it is applied on a plant. Naturally occurring cytokinins may well behave differently though, as there is some evidence that cytokinins are synthesized in roots and translocated up into the shoot system (p. 290). Certainly it is known that some synthetic cytokinins are translocated quite readily in plant tissues.

Auxins are not the only factors concerned in root formation. A supply of sugar is necessary, as well as other nutrients. The stimulating effect of leaves on initiation in stem cuttings

may be due in part to their production of nutrients, and perhaps also to other hormonal substances more specific in promoting root formation in conjunction with auxin.

Lateral Root Initiation

In contrast to the fairly ready regeneration of roots in shoot cuttings, excised roots of most species growing in sterile culture normally form only further root tissues, including the initiation of lateral roots, and only relatively rarely are shoot buds initiated. Auxin is in some way involved in the formation of lateral roots as well as of adventitious roots. Immersion of the main root of a dicotyledonous seedling in a solution of an auxin results in a reduction of main root extension but a stimulation of lateral root initiation. The subsequent growth of the newly produced lateral roots is also inhibited by the auxin solution. Thus auxins, at other than very low concentrations, stimulate the formation of roots but inhibit their subsequent elongation. The result is that a root immersed in a solution of auxin becomes stunted, and possesses rows of newly emerged but suppressed lateral roots.

GENERAL ASPECTS OF REGENERATION

Sinnot has defined regeneration as "the tendency shown by a developing organism to restore any part of it which has been removed or physiologically isolated and thus to produce a complete whole". This broad definition includes a wide variety of phenomena, but we can distinguish a number of general aspects of regeneration. Firstly, we have seen that we can apply the term to the initiation of shoot and root meristems in a disorganized mass of callus, which may be growing in aseptic culture or may form at the surface of a cutting in response to wounding. This is a remarkable phenomenon, even though it may be so familiar that we come to accept it as commonplace. We have no conception as to the nature of the factors operating whereby in a mass of disorganized callus a high degree of organization emerges, but we have already suggested (p. 39), that the apical meristem of the shoot or root is a stable configuration which, as it were, "crystallizes" out under certain conditions.

It is important to realize that in regeneration we see in operation the processes which determine normal development. When the normal course of development is disturbed by wounding or in other ways, compensating events occur which tend to restore the normal situation. Thus, it would seem that the normal form of the plant represents an equilibrium state, and that when this equilibrium is disturbed, built-in control mechanisms operate to restore the equilibrium. This phenomenon is well illustrated in the regenerative properties of shoot and root meristems. We have already seen that if a shoot apex of *Lupinus* is divided by two vertical cuts at right angles, then each segment of the original apex is able to regenerate into a normal apex (p. 39). Similar experiments have been successfully carried

out with root apices. Further examples are seen in the regeneration of vascular tissue (p. 118) and in the formation of a phellogen when the surface of a stem is cut or damaged.

So far we have discussed the problems presented by the spontaneous development of organized meristems within unorganized meristematic tissue. A further problem concerns the resumption of cell division in previously differentiated, non-dividing cells, which follows wounding. In some cases regeneration of root primordia takes place in callus tissue which has developed at the cut basal surface of a shoot cutting, while in other cases the root primordia may be formed by the resumption of cell division in stem tissues, such as the pericycle. In either case, however, it is clear that the isolation of a piece of stem or other organ results in renewed cell division, and the question arises as to what causes this cell division. There is some evidence that the wounding of plant tissues results in the release of "wound hormones", which stimulate cell division. Whether such substances are involved in all cases of cell division following wounding is not clear. In any case, however, it is clear that certain differentiated cells of the stem or root become "dedifferentiated" when they resume meristematic activity.

The phenomenon of regeneration provides strong evidence that the process of differentiation in many types of plant cell does not involve any loss in their genetic potentialities, so that they remain "totipotent".

Although the totipotent behaviour of individual cells of a number of plant species has been demonstrated experimentally (Fig. 6.5), it is nevertheless wise to be cautious in assuming that all living, nucleated, plant cells are totipotent. Until regeneration of whole plants has been seen to occur from isolated cells of all types known to occur in the plant body (e.g. parenchyma, palisade and spongy mesophyll cells, companion cells), the case cannot be regarded as proven. Even so, as we have mentioned earlier (p. 148), plants have been regenerated from adventive embryos formed from cells derived from the root, hypocotyl, stem, petiole, embryo, and even pollen. Cells from the lamina of green leaves do appear to be more recalcitrant in demonstrating their presumed totipotentiality. A number of workers in the field of plant cell and tissue culture, particularly Steward, nevertheless consider that any free cell from a higher plant will, if provided with the right stimuli (nutritional and hormonal), regenerate a whole new plant taking one of the alternative routes toward organization described above.

It seems clear that the developmental activities of most callus cells are restricted in some way, and that further restraints are imposed should differentiation of vascular tissues, shoot-buds and root initials occur. Thus, the cells of undifferentiated callus are usually capable of unlimited division, but if a bud is regenerated, then the cells which become part of the leaf primordia are subject to considerable restraints with respect to the planes of their divisions, and they are no longer capable of unlimited division so long as they remain part of the leaf. We do not know how a cell becomes restricted when it is a part of a tissue system, but possibly some control over each cell is exerted by its neighbours through the plasmodesmata, which connect the protoplasts of adjacent cells.

The discovery that the regeneration of buds and roots by callus tissues can be regulated by the relative concentrations of auxin and cytokinin in the culture medium has led to the

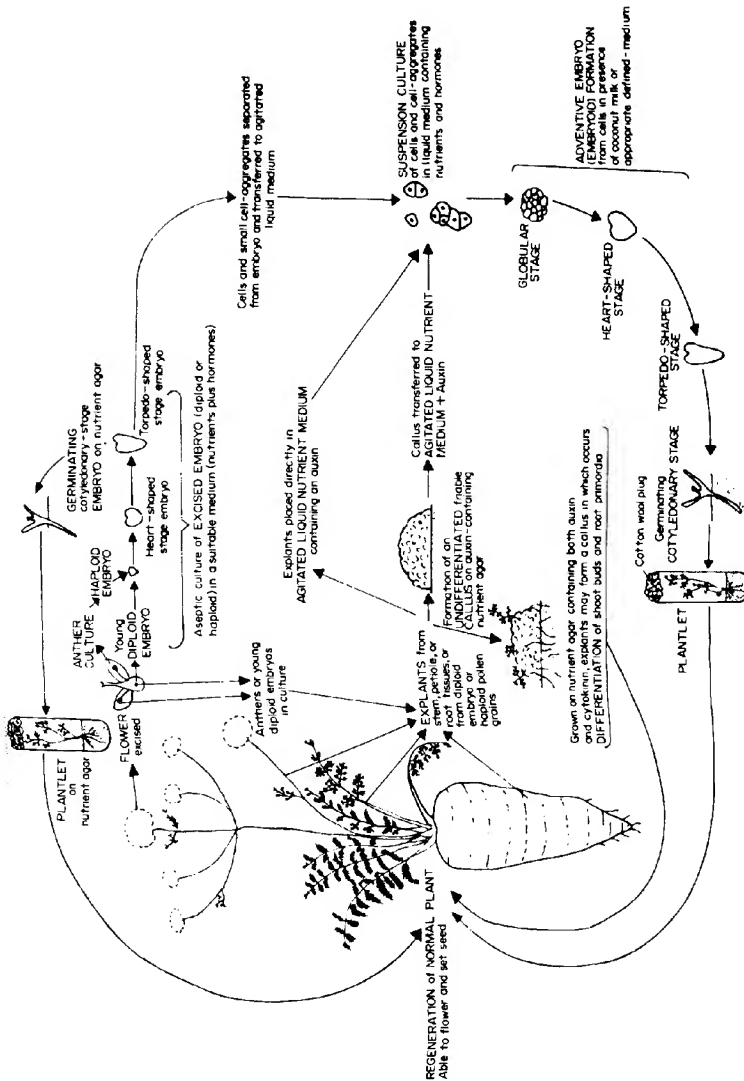


FIG. 6.5. Diagram to illustrate alternative pathways of regeneration of a whole new plant from tissue explants derived from various parts of a carrot plant. The regeneration cycles can be repeated indefinitely. (Adapted, with additions, from F. C. Steward *et al.*, *Brookhaven Symp. Biol.* 16, 73, 1963.)

suggestion that these hormones play an important role in "organ formation". However, it should be noted that the primary effect of the hormones is upon the initiation of shoot and root apical meristems, and we do not find callus cultures giving rise directly to organs of determinate growth, such as stems and leaves, although these may be formed as a result of the subsequent growth of the meristems. Unless we are prepared to call a shoot meristem an "organ", which seems inappropriate, it is not strictly accurate to say that cytokinins promote organ formation, although it is perhaps more justifiable to say this of the initiation of root primordia in response to auxin.

Nevertheless, the initiation of two kinds of apical meristem in response to different hormone levels is a highly interesting effect and raises the question as to whether differences in the levels of endogenous auxins and cytokinins in different parts of the plant may play a role in normal development. The observation that there is an increase in the levels of endogenous cytokinins in the apical ends, and of auxins at the basal ends of cuttings of chicory root (p. 156) and that these changes precede the initiation of adventitious buds and roots, suggests that these hormones are important in natural regeneration. We shall return to a discussion of the possible role of growth hormones in morphogenesis in Chapter 13.

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CHAPTER 7

Growth Responses to Directional Light and to Gravity

Introduction—The Environmental Control of Development

So far, we have considered mainly the internal processes which are involved in and may control growth and differentiation in plants. However, land plants are subject to a wide variety of external influences, including light, temperature, water stress, gravity and so on, which may modify the growth and development of the plant to a greater or a lesser degree. Some of these external influences may have deleterious effects upon the plant and may constitute a hazard, as with freezing temperature or drought. Animals also have to survive under these conditions but they are frequently able to *avoid* the stresses by migration or hibernation, whereas land plants, being sedentary, have little scope to take avoiding action and must *tolerate* the environmental stresses if they are to survive.

However, apart from the direct effects of environmental stress on plant tissues, variations in certain other external factors may be advantageous to the plant by providing it with information about the environment which facilitates adaptation of the plant to its local situation. Thus, gravity never constitutes an environmental stress, but the capacity of the plant to detect the direction of gravity provides information whereby it can regulate its orientation in relation to the soil surface. In such cases the environmental factor acts as a *signal* or *stimulus*, and the plant responds in an active, "programmed" manner, which enables it to become better adapted to its environment. Thus, it is useful to distinguish between:

- (1) effects of environmental factors which are direct and non-adaptive (e.g., freezing injury); and
- (2) adaptive, programmed responses, in which the environmental factor acts as a signal or stimulus as in geotropism.

Adaptive responses to the environment are numerous and varied and we shall be considering a number of these in the remaining parts of the book. In the present chapter we

shall confine ourselves to a consideration of growth curvatures that occur in response to directional light stimuli and to gravity.

GROWTH MOVEMENTS

It is one of the characteristic properties of living organisms that they have the ability to perceive and respond to changes in external or internal conditions. The change in external conditions is called the *stimulus* and the resulting change in the plant is called the *response*. The response may take place in various ways, but very frequently stimulation results in *movement*.

All plants have the capability of movement. In lower plants, such as many algae, fungi and bacteria, movement of the whole organism occurs. In higher plants the capability of movement is restricted to individual organs or parts of the whole organism. The process of straight extension growth itself can perhaps be regarded as such a movement, in that, for example, the root tip moves through the soil as a result of growth. Other types of movement result from *differential growth rates*; that is, the organ shows different rates of growth on opposite sides and this results in the bending of the organ in one direction. Such movements are termed *growth movements*. Not all plant movements are growth movements. Some are brought about by reversible turgor changes in tissues at the base of each leaf and there is often a specialized structure, the pulvinus, at which bending occurs due to reversible changes in turgor. (A pulvinus is the swollen base of a leaf or leaflet which contains a high proportion of thin-walled parenchyma.)

We have already mentioned (Chapter 3) one growth movement, that of phototropism, where movement of an organ occurs in response to unilateral illumination. A tropic movement of this type is thus a response to an external stimulus. In the case of phototropism the stimulus is light, but other external stimuli such as gravity, water, chemicals, heat or mechanical contact can also induce growth movements.

Where the direction of the response is related to the direction of the stimulus, we speak of a *tropic* response, or *tropism*; but in many cases the direction of movement does not bear a direct relation to that of the stimulus, and we then speak of a *nastic* response. Thus, a curvature of a shoot towards the more illuminated side is a phototropic curvature, while the opening or closing of flowers with a change of light intensity all round is a *photonastic* one. All tropic responses are induced by directional or unilateral stimuli, whereas in nastic responses the stimulus may be diffuse. Another example of a nastic growth response is seen in the opening and closing of certain flowers, such as those of crocus, in response to changes in temperature (thermonasty). When the temperature rises growth is faster on the inner side of the petals, so that the flower opens, whereas the reverse is true when the temperature falls. Not all nastic movements are growth movements, some being brought about by reversible turgor changes in pulvini, as in the movements of French bean (*Phaseolus vulgaris*) leaves in response to light and dark.

We shall consider phototropism and geotropism in more detail below, but a few

examples of other types of response may be mentioned here. The tropic movement of a plant organ in response to an external chemical stimulus is named *chemotropism*. An example is seen in pollen-tube growth down the style towards the ovules, for the directional stimulus for the elongating pollen tube is provided by certain chemicals present in the ovule and ovary wall, although their precise nature is not known. The familiar sight of tendrils of a pea plant, or other climbing plants, twining around a support is a good example of a response to contact or mechanical stimulus, and is called *haptotropism* (or *thigmotropism*). Many roots respond to differences in soil moisture content and grow towards regions of wetter soil (*hydrotropism*). There are many other examples of both tropic and nastic responses which, for reasons of space, we cannot describe further here.

A few growth movements in plants do not obviously result from any external stimulus, but appear to be spontaneous movements arising from causes within the plant itself. Examples are straight extension growth, nutations and epinastic movements. A *nutational movement* is seen in the growth of a plant stem, for it does not grow straight upwards, but performs a series of rhythmic movements which result in the shoot tip oscillating about the longitudinal axis. This particular type of nutation, called *circumnutation*, is very pronounced in both shoots and tendrils of climbing plants, and may confer a biological advantage in the finding of a support. An example of an *epinastic movement* is seen in the petioles of many species, which show a growth curvature, the upper side growing more rapidly than the lower, resulting in a downward movement of the lamina as the leaf grows. This downward movement of leaves with increasing age is known to result from an internal stimulus, and not from an external one such as gravity.

PHOTOTROPISM

Phototropism is the term applied to the phenomenon whereby a plant organ responds to a directional ("unilateral") light stimulus by undergoing a directional, or differential, growth response. As we saw earlier (p. 48), it was studies of phototropism which led to the discovery of auxins. Phototropism differs from photomorphogenesis (Chapter 8), in that photomorphogenic responses are neither dependent on a *directional* light stimulus nor do they show characteristics that are related to the direction from which the photomorphogenic light stimulus may be received.

In general, stems and other aerial portions of plants are positively phototropic (i.e. they bend towards the light source), while roots and other underground organs are negatively phototropic (they bend away from the light source). There are, however, many exceptions to these rules; for example, some tendrils and stems are negatively phototropic, and many roots non-phototropic or even positively phototropic when young, becoming negatively phototropic only later on. There is no doubt, nevertheless, that phototropism is of great importance in determining the direction in which plant organs develop under natural conditions. Phototropic responses, by definition, can occur only in those parts of a plant

that retain the capacity for growth, particularly elongation growth. Thus, one sees phototropism in the young growing stems, leaves and roots of higher plants, and also in the sporangiophores of some fungi, in the sporophores of mosses, and chloronemata of ferns. The majority of studies of phototropism have, however, been made on etiolated coleoptiles of grass seedlings (particularly of oat, wheat, maize, and barley) and on fungal sporangiophores. The reasons for this are the convenience which such organs offer in experimentation, and their great sensitivity to directional illumination. Although such work has yielded much valuable information on phototropism, one cannot be confident that the derived concepts are directly applicable to other organs, such as green leafy shoots, growing under natural daylight conditions.

As in all environmentally induced adaptive responses, phototropism involves first of all *perception* of the stimulus (directional light) which is followed by development of the *response* (directional growth). One can discuss separately the mechanisms of perception and response, but keeping in mind that the two processes are linked together in the plant.

The earliest major advance in our understanding of phototropism was taken by Charles Darwin (1880) in his studies of the phototropic responses of coleoptiles (Chapter 3). In particular, he demonstrated that the tip of the coleoptile was the region of perception of a directional light stimulus, and that the differential growth response occurred lower down. Although it has subsequently been found that more basal parts of coleoptiles may also show some sensitivity to directional light, Darwin's basic results and conclusions have been amply confirmed. Thus, the coleoptile tip is the region of maximum photosensitivity, and following receipt of a directional light stimulus it transmits basipetally some influence which elicits differential growth in the elongating parts of the coleoptile.

The Nature of the Phototropism Photoreceptor

(a) *Dose-response relationships.* Early in the twentieth century, before the discovery of auxin, experimental work on phototropism concentrated upon biophysical aspects. As early as 1909 it was established by Blaauw that in both grass coleoptiles and *Phycomyces* sporangiophores, the Bunsen-Roscoe Reciprocity Law holds over a rather wide range of light intensity and time. The reciprocity law states that when only a single photoreceptor is operating, then the photochemical effect of light remains the same if the *quantity*, or dose, of light (i.e. intensity \times time of irradiation) remains the same. Blaauw's observations therefore indicated that a single photoreceptor is operative in phototropism, and led logically to investigations of action-spectra for phototropism with a view to identifying the photoreceptive pigment concerned in the perception of directional light.

The dose-response relationships for phototropism are, nevertheless, much more complex than first appeared the case. Thus, in the case of etiolated coleoptiles it has been found that with increase in the quantity of stimulus there is an increase in the bending response towards the light source until a maximum is reached (with approximately 0.1 J m^{-2} light energy), above which, with increasing quantity of stimulus, the response falls off until at a certain

value the initial "positive" curvature may even be reversed and a "negative" curvature (i.e. away from the illuminated side) occur. With still greater quantities of light stimulus the curvature may again become positive. The three regions of the dose-response curve are called System I (or "first-positive"), System II ("first-negative"), and System III ("second-positive") curvatures (Fig. 7.1). The Bunsen-Roscoe Reciprocity Law has been found to be valid only for System I and System II responses, and the System III (second-positive) curvature response does not obey the law. Because of this, an accurate action spectrum can be determined only for the responses to lower light dosages (ie. those eliciting System I or System II curvatures), and there remains a possibility that System III curvatures involve a different pigment, or pigment system, from that concerned in Systems I and II. However, most workers in this field consider that it is probable that the same photoreceptor operates in all types of phototropic response, and that deviations from the reciprocity law result from as yet undiscovered interactions between the pigment and other cell components. However, it must again be emphasized that most studies of phototropism have been conducted on the first-positive (System I) responses of etiolated coleoptiles, and it is not possible to say how relevant are the results to green plants growing under normal daylight conditions.

(b) *Action spectrum studies.* Research to identify the phototropic photoreceptor has

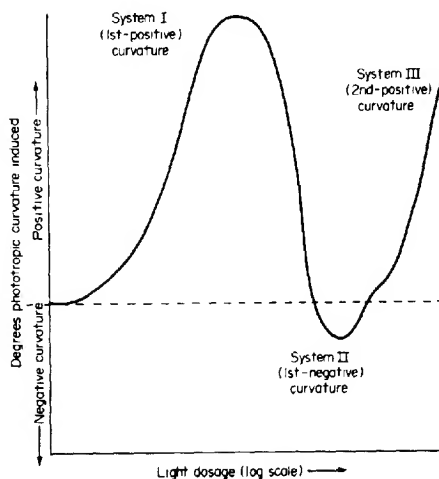


FIG. 7.1. Summary of Systems I, II and III phototropic curvatures of oat (*Avena*) coleoptiles exposed to varying dosages of monochromatic blue light. Coleoptiles of other cereals show a similarly shaped curve, except that System II may not be clearly negative. (Adapted from W. R. Briggs, in *Photophysiology*, Vol. 1 (ed. A. C. Giese), pp. 223-71, Academic Press, New York and London, 1964.)

naturally centred around attempts to match the action spectrum for phototropism with the absorption spectra of likely photoreceptor pigments. Results obtained do not yet permit a definite decision to be taken on the identity of the pigment. The action spectrum for the first-positive curvature in coleoptiles (Fig. 7.2) shows that maximum curvature occurs in response to light of the blue wavelengths (maxima at about 445 and 474 nm, and a shoulder in the 425 nm region). Another, much lower, maximum occurs near 370 nm (the near

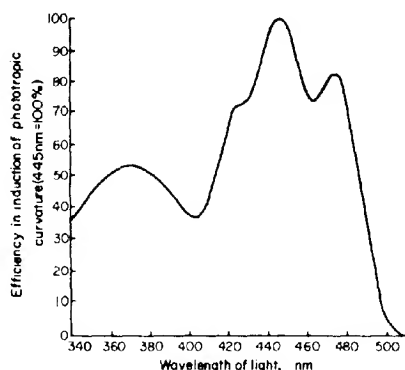


FIG. 7.2. Action spectrum for first-positive (System I) phototropic curvature in the *Avena* coleoptile. Maximum curvature occurs with unilateral light of the blue wavelengths. (From K. V. Thimann and G. M. Curry, *Comparative Biochem.* **1**, 243-306, Academic Press, 1960.)

ultra-violet). There is a sharp cut off in activity at about 500 nm with no activity at wavelengths longer than this. Such an action spectrum suggests that the phototropism photoreceptor is a yellow pigment, and detailed action spectrum studies on phototropism in coleoptiles have narrowed down possible photoreceptors to just a few yellow substances known to occur in plants. At present the two strongest possibilities are a carotenoid or a flavin pigment. The absorption spectra of β -carotene and riboflavin are shown in Fig. 7.3, and it can be seen that although both have absorption maxima in the blue wavelengths, neither of the absorption spectra are good matches with the action spectrum for phototropism (Fig. 7.2). The carotenoids do not have a peak of absorption at or near 370 nm, and flavins lack the complex pattern of absorption in the blue region. On the other hand, the carotenoid absorption pattern in the blue region matches well the action spectrum, and flavins do have a strong absorption peak at 370 nm. Both flavins and carotenoids are present in greatest concentrations in the apical parts of coleoptiles, with much lower levels in more basal, less light-sensitive, regions.

Thus, action spectrum studies in relation to naturally occurring pigments such as carotenoids and flavins have not produced a definite answer on the question of the identity of the

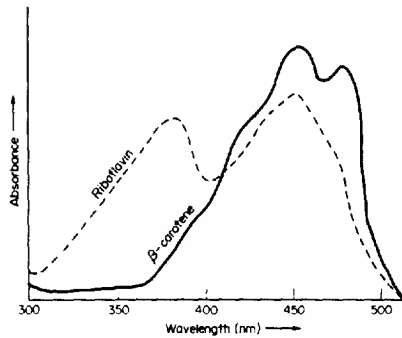


FIG. 7.3. Absorption spectra for riboflavin and β -carotene. Neither spectrum matches the action-spectrum for phototropism (see Fig. 7.2), but when the determination on riboflavin is made in a lipoidal solvent then its absorption spectrum more nearly resembles the action-spectrum.

phototropism receptor. One of the difficulties in this type of investigation is that the absorption spectrum of a compound can be greatly affected by the particular solvent in which it is dissolved during spectrophotometry. For example, it has been observed that the absorption spectrum for riboflavin as shown in Fig. 7.3 is altered so as to much more closely match the phototropism action spectrum (Fig. 7.2) when the pigment is dissolved in a suitable lipoidal solvent. Because the absorption spectrum of a pigment is so susceptible to chemical environment, it is really necessary to measure it *in vivo*, or at least in the medium in which it occurs naturally, to be able to make a meaningful comparison with the action spectrum for phototropism. This has not proved possible to the present time.

Other possible reasons that have been proposed to explain the lack of match between the phototropism action spectrum on the one hand, and measured absorption spectra of individual plant pigments on the other, are: (a) that the active pigment may exist in the plant in a complex with another substance or substances, (b) that it may be affected by "masking" by other light-absorbing substances, and (c) that a carotenoid and flavin co-operate in light absorption to act as the phototropism receptor, but this would not, of course, be expected from the dose-response studies which in the main have indicated that a single photoreceptor is involved.

Recent research is, nevertheless, leading to the view that the phototropism photoreceptor may indeed be a complex between a flavin (perhaps riboflavin) and a b-type cytochrome, and that this flavin/b-cytochrome complex is membrane-bound. The work which initiated this currently attractive hypothesis was first conducted in 1974 and 1975 on the cellular slime mould *Dictyostelium discoideum* and on the fungi *Phycomyces blakesleeana* and *Neurospora crassa*. These studies were not on the phenomenon of phototropism, but the results have attracted the interest of other workers in relation to phototropism in coleoptiles. The

reason for this is that it was found that blue light (maximum approximately at 465 nm) brings about a photoreduction of b-type cytochrome in these lower organisms, which is a region of radiation *not* absorbed to any significant extent by cytochrome itself. Further work indicated that a flavin absorbs the blue light, and close association between the flavin and cytochrome results in the latter being photoreduced. Preliminary cell-fractionation and other studies on *Neurospora* and on *Zea mays* coleoptiles have suggested that a similar flavin/b-cytochrome photoreceptor is present in the cells of both organisms and is associated with the plasma membrane. Only further detailed research can tell us whether this flavin/cytochrome complex has significance in the understanding of phototropism.

Phytochrome, the red and far-red absorbing photoreceptor concerned in the numerous photomorphogenic phenomena in plants (Chapter 8), does not appear to be directly concerned in phototropism. However, it is known that exposure to red-light (non-directional) does affect the sensitivity of plant organs to directional blue light, and the effect of red can be reversed by exposure to far-red light. Such results indicate that phytochrome is the pigment which mediates red light effects in phototropism, but it is not known how phytochrome is linked to the phototropism system.

The Mechanism of Redistribution of Growth in Phototropism

We have seen earlier (Fig. 3.1) that various early experiments led to the conclusion that there is a transmission of auxin from the coleoptile tip to the basal tissues of the shaded side. Over the years, several theories of phototropism involving auxin as a correlation factor have been proposed. These are:

- (1) The *Cholodny-Went theory*, which was put forward independently by Cholodny and Went in the 1920s. This theory suggests that a phototropic stimulus induces a *lateral translocation of auxin* across the photosensitive region of a coleoptile, leading to a higher auxin concentration in the darker half which consequently elongates more rapidly than the illuminated side—i.e. a positive phototropic curvature ensues.
- (2) A theory invoking *photodestruction of auxin* in the illuminated tissues (see also p. 52), which produces a differential in auxin concentration between illuminated and darkened regions.
- (3) The suggestion that the rate of auxin *synthesis* is lower in the illuminated than in the darker parts of the coleoptile tip.

Apart from these three theories, there is a fourth concept, that of a *light-growth reaction* not necessarily involving auxin. This suggests that when light impinges on a plant cell it affects its growth rate, either stimulating or suppressing it, and is considered later in this chapter (p. 172).

The original evidence in favour of the Cholodny-Went theory was based upon work by Went in 1928, who made measurements of the quantities of auxin which could be collected in agar blocks placed below the illuminated and shaded sides of cut coleoptile tips (Fig. 7.4).

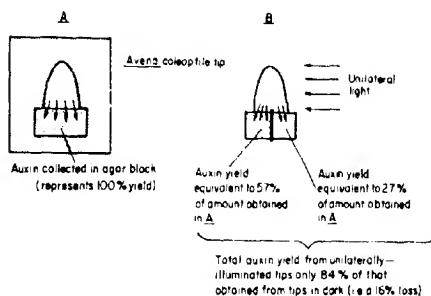


FIG. 7.4. Went's original experiment demonstrating that an oat coleoptile tip, when exposed to unilateral light, transmits more auxin to an agar block below the shaded side than to the block below the illuminated side (B). The total auxin yield from illuminated tips was 16 per cent less than that from tips maintained in darkness (A). This apparent loss was probably not significant (see text). (From F. W. Went, *Rev. Trav. Bot. Neerl.* 25, 1-116, 1928.)

It was found that unilateral light increased the proportion of total auxin in the agar blocks positioned below the shaded side. Went found that 57 per cent of the total auxin that had been obtained from the non-illuminated tip was collected from the shaded half of a unilaterally illuminated tip, and only 27 per cent from the illuminated half (Fig. 7.4)—i.e. revealing a 16 per cent overall fall in total auxin yield following illumination, the remaining auxin being preferentially distributed to the darker region. The 16 per cent fall in auxin resulting from exposure to light was regarded as insignificant by Went, who concluded that a unilateral light stimulus induces lateral migration of newly synthesized auxin towards the shaded side.

In his experiments, Went used a unilateral light dosage which would be expected to induce a first-positive curvature, but a number of other workers have repeated Went's experiments using light dosages which would induce first-negative or second-positive curvatures. In all these cases an auxin differential was obtained, with the greatest amount of auxin emerging from the basal cut surface of a coleoptile tip corresponding to the most rapidly elongated side of the responding coleoptile.

Although Went regarded the 16 per cent loss in total auxin which occurred on phototropic stimulation as being insignificant, a number of other workers considered that photodestruction of auxin in the illuminated tissues may play an important part in phototropism. It was found by Galston and co-workers that blue light can be absorbed by natural plant pigments such as riboflavin, and the absorbed energy utilized in the photo-oxidation of IAA (see also p. 52). Similarly, work by Zenk has shown that a naturally occurring xanthophyll, violaxanthin, absorbs light of wavelength of 450 nm which can energize the destruction of the auxin, naphthalene-acetic acid. Nevertheless, Briggs and his associates, and Gillespie and Thimann have shown that light of the dosage necessary to produce phototropic curvatures does *not* decrease the overall auxin levels in coleoptile tips. There is little

justification, therefore, for regarding photodestruction of auxin as playing any important part in phototropism in coleoptiles.

Even so, it is obviously of critical importance to know whether there was any real significance in the 16 per cent loss in auxin following light stimulation in Went's original experiment. Because of this, Briggs and his co-workers in California carried out a series of experiments in 1957 using corn coleoptiles (*Zea mays*), which essentially confirmed earlier work by Boysen-Jensen in that the results showed that the lateral differential in auxin concentration occurring in phototropically stimulated tips could be a consequence only of light-induced lateral transport of auxin from the light to the dark side. Briggs collected in agar blocks the auxin which diffused from coleoptile tips which had been subjected to various treatments, and then determined the quantity of auxin present in the blocks by means of the Went *Avena* curvature test. The amount of auxin produced by the tips kept in total darkness was similar to that produced by illuminated tips, whether or not the tips had been completely separated into two vertical halves by a thin piece of glass (Fig. 7.5a-d). This result argues against the hypothesis of photodestruction of auxin and photoinhibition of auxin synthesis. Further, it was found that when a tip was stood on two separate agar blocks and only partially bisected at its base by a glass sheet, and was exposed to unilateral light incident at right angles to the glass plate, then significantly more auxin diffused into

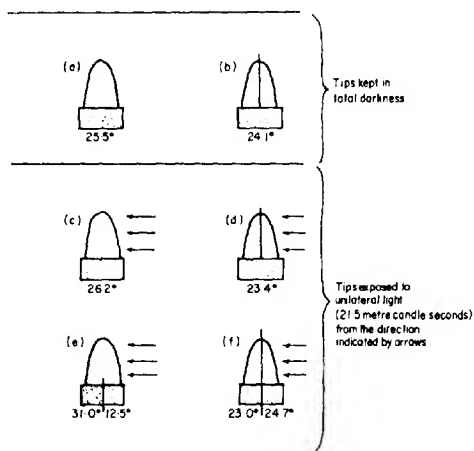


FIG. 7.5. Auxin diffusion into agar blocks from variously treated coleoptile tips of *Zea mays*. The figure under each agar block indicates the degrees curvature produced by that block in the Went *Avena* curvature test for auxin. The vertical line running through some of the tips and agar blocks represent an impervious glass barrier. Twice as much auxin was obtained in (e) and (f) than in (a)-(d), because each agar block had been in contact with six half-tips, the equivalent of the three whole tips placed on the agar blocks in (a) to (d) inclusive. (From W. R. Briggs, R. D. Tocher and J. F. Wilson, *Science*, **126**, 210-12, 1957.)

the agar block below the half of the tip remote from the light source (Fig. 7.5e). When, however, the whole of a coleoptile tip was bisected and separated by glass and exposed to the same conditions as a partially separated tip, then it was found that there was no difference in the amounts of auxin diffusing into the separate agar blocks below the "light" and "dark" halves (Fig. 7.5f). If either auxin destruction or inhibition of synthesis was responsible for the observed differential in the partially split tips one would have expected that a total glass barrier, as in Fig. 7.5f, would make no difference to the differential distribution of auxin observed in the partially split tips. The fact that a total glass barrier completely prevented the establishment of an unequal distribution of auxin led Briggs to conclude that the observed differential distribution of auxin that occurs in phototropically responding coleoptile tips is a consequence of lateral movement of auxin towards the dark side, and is not a result of photodestruction of auxin or of photoinhibition of synthesis.

In summary, most available evidence indicates that when a coleoptile tip is illuminated from one side, there is first of all *perception* of the light-stimulus, followed by *transverse migration* of endogenous auxin molecules within the tip. In the case of positive phototropic responses auxin moves towards the darker side, which in turn means that more auxin is transmitted to the region of response in the coleoptile below the darker half of the tip (Table 7.1), resulting in greater elongation growth of that region and consequent bending of the whole coleoptile towards the light source. There is some evidence that negative phototropism in coleoptiles similarly involves transverse migration of endogenous auxin, but in this case towards the light source so that the illuminated side contains greater amounts of auxin (Table 7.1) and elongates more rapidly than the darker side.

TABLE 7.1. Effects of varying unilateral light dosages upon direction of phototropic curvature in oat coleoptiles and amounts of auxin diffusing from illuminated and shaded halves of the tips. Auxin quantities measured in Went *Avena* curvature test. (From M. Wilden, *Planta*, 30, 286-8, 1939-40)

Light dosage (mc s)	Direction of curvature	Ratio of auxin diffusing from tip halves (illuminated:shaded)
1,500	positive	17:83
11,400	negative	62:38
150,000	positive	36:64

The concept of lateral migration, or transport, of auxin as a major component of the phototropic response mechanism has also received support from results of experiments that have demonstrated lateral migration of radioactively labelled exogenous auxin (^{14}C -IAA, or ^3H -IAA) in response to unilateral illumination of coleoptiles. Especially convincing have been experiments by M. B. Wilkins and associates, in which ^3H -IAA of high specific activity was applied by micropipette to the apical end of *intact* coleoptiles (i.e.

the radioactive auxin was administered at extremely low concentration to the most photosensitive region of the organ).

Although we may assume that transverse migration of auxin occurs under the influence of directional blue light, the mechanism by which this occurs is unknown. An attractive possibility that was emphasized during the 1930s and 1940s was that an electrical potential difference between light and dark sides of a unilaterally illuminated organ is involved. Such bioelectrical potentials of approximately 100 mV occur spontaneously in phototropically responding organs, and it was suggested that ionized acidic auxin molecules migrate electrophoretically towards the more electrically positive dark side. But more recent research with refined instruments has shown that the lateral potential is not set up until after lateral auxin transport has already started. Furthermore, decapitated, auxin-depleted, coleoptiles do not develop such a transverse potential unless exogenous auxin is administered. It seems, therefore, that the transverse potential is a consequence, rather than cause, of differential growth rates in the dark and light halves of a phototropically responding coleoptile.

The Light-growth Reaction

In the early years of this century it was noticed by Blaauw that in both grass coleoptiles and *Phycomyces* sporangiophores, blue light administered symmetrically results in transient changes in growth rates. This has been called the "light-growth reaction". Blaauw and subsequent workers have established that in *Phycomyces* there is first an elevation of growth rate, then a depression followed by a return to the normal rate. In coleoptiles there occurs at first (3–20 minutes blue light) a depression, then (25–45 minutes) an elevation, and then a return to the control rate after about an hour. The action spectrum for these light-growth reactions matches the action spectrum for phototropism for the particular organism. Some workers, including Blaauw, concluded that phototropism can be explained completely in terms of the light-growth reaction.

Thus, it has been sometimes envisaged that in the oat coleoptile, where blue light initially suppresses growth, a unilateral irradiation with blue light would set up the type of growth asymmetry required for commencement of a positive curvature (i.e. towards the light source). However, it is difficult to understand what would happen after the first 20 minutes, when the transitory growth inhibition is converted to a transitory increase in growth rate, but it is possible that the time relationships are different under conditions of unilateral irradiation.

In the case of the light-growth reaction of the *Phycomyces* sporangiophore, illumination results in an initial increase in growth rate, which at first sight makes it difficult to see how it can explain curvature toward a unilateral light source. However, elegant experiments have revealed that the reason is that the sporangiophore acts as a cylindrical lens, focusing light internally onto the "shaded" side and presumably causing there a greater photochemical reaction and greater acceleration of growth on that side. Immersion of the

sporangiophore under paraffin oil results in it executing a negative rather than the normal positive phototropic curvature, presumably because the high refractive index of the oil disperses incoming light and prevents it being focused on the far side of the sporangiophore; thus, the light-growth reaction would take place to a greater extent on the illuminated side, and curvature away from the light source would take place. Phototropism in the *Phycomyces* sporangiophore does, therefore, appear to result from a blue light-growth reaction with the lens effect being essential for normal positive curvatures. On the other hand, the involvement of a light-growth reaction in phototropic responses of coleoptiles or other higher plant organs has never been convincingly demonstrated. Nevertheless, and inexplicably at present, it has been reported that immersion of oat coleoptiles in paraffin oil reverses the direction of phototropic curvature just as in *Phycomyces* sporangiophores. Also, positive phototropism in chloronemata of ferns appears to involve a light-growth reaction, but in this case it is red, and not blue, light which is effective, and it is therefore likely that phytochrome is involved (Chapter 8).

Phototropism in Green Plants

The great majority of experiments so far conducted on phototropism have been concerned with the behaviour of etiolated organs, particularly coleoptiles. It is not possible to say how relevant the findings and derived concepts are to the situation in green leafy plants. Very few experiments indeed have been done on light-grown plants—either monocotyledonous or dicotyledonous, but available evidence indicates that light-grown plants are much less sensitive to directional blue light than are etiolated coleoptiles, responding only to energies corresponding to the second-positive response of etiolated coleoptiles. Nothing is known of the photoreceptor pigment(s) in phototropism of green plants. An added complication in leafy shoots, particularly in dicotyledonous plants, is that it has been found that most of the auxin required for elongation of the stem comes from the young expanding leaves near the apex. In other words, the lamina of a young leaf exports auxin via the petiole into the stem. Consequently, any phototropic curvature of such a stem must be a result of some alteration in the distribution, or quantity, of auxin coming from the leaves. In the case of sunflower plants (*Helianthus annuus*), the leaves are arranged in pairs on opposite sides of the stem (decussate arrangement) and it has been found that each leaf of a pair will, if both are illuminated to the same degree, supply equal quantities of auxin to the stem (Fig. 7.6A). If, however, one leaf of a pair is more brightly illuminated than the other because of its orientation in relation to the incident light (Fig. 7.8B), then the leaf receiving a higher intensity of light produces a greater quantity of auxin than its partner. This perhaps results in the side of the stem beneath the brightly illuminated leaf receiving more auxin and consequently growing at a more rapid rate than the other side, causing the stem to execute a positive phototropic curvature, until the position is reached whereby both leaves receive light at equal angles of incidence (Fig. 7.6C).

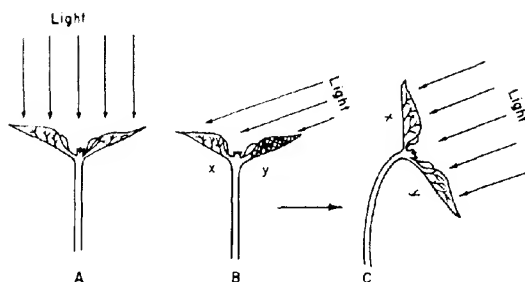


FIG. 7.6. Effect of light intensity on auxin production in young leaves of *Helianthus annuus*, and on the induction of a phototropic curvature in the stem. A. Both leaves illuminated equally and producing equal quantities of auxin. B. The illumination of leaf X is higher than of leaf Y, because of the difference between the angles at which the light strikes them. Hence there is greater auxin production by leaf X than by leaf Y. C. Phototropic curvature resulting from unequal auxin production by the opposite leaves. (From H. Shibaoka and T. Yamaki, *Sci. Papers Coll. Gen. Educ. Tokyo*, 9, 105-26, 1959.)

GEOTROPISM

This term is applied to growth movements induced by a gravitational stimulus. Growth of an organ towards the centre of the earth is termed *positive geotropism*, and growth away from the centre of the earth *negative geotropism*. Positively and negatively geotropic organs, such as the stem and root of the main plant axis, which align themselves parallel to the direction of the gravitational pull, are said to be *orthogeotropic*. When the axis of an organ come to lie at right angles to the direction of the gravitational field it is said to be *diageotropic* (e.g., rhizomes of Solomon's seal, couch grass, etc., or stolons of potato and strawberry). Where an organ becomes oriented at intermediate angles (i.e. between 0° and 90° , or between 90° and 180° from the vertical), is it said to be *plagiogeotropic* (e.g. lateral branches are very often so). Most main roots are positively geotropic. The rhizomes of many mosses are sometimes positively geotropic, but, in general, positive geotropism is not well developed in lower plants. Negative geotropism is shown by the stems of higher plants, by the sporangiophores and sporophores of many fungi, and by the foliage shoots of mosses. While many rhizomes and stolons are diageotropic, lateral stems and lateral roots of the first order and foliage leaves are commonly plagiogeotropic. Lateral shoots and lateral roots of a higher order generally possess little geotropic sensitivity and are consequently said to be *ageotropic*.

Moving a plant from its usual vertical position to a horizontal one causes gravity to act across the width of stem and root. This results in growth curvature responses, whereby the stem bends to grow upwards and the root bends to grow downwards. This can easily be demonstrated with a young seedling, such as that of *Zea mays* or mustard.

Geotropism resembles phototropism in a number of respects. It is a response to a directional stimulus; it involves differential elongation growth leading to curvature; perception of the gravitational stimulus occurs in the apical region of coleoptile, shoot or root; and there is transmission of a growth-regulating factor from the region of perception to the region of response.

Gravity Perception

Some sort of *geoperceptive* mechanism must exist in plants, which senses the direction of gravity in relation to the orientation of the organ. It has been known for many years that the geotropic response is a "threshold" phenomenon, in that a gravitational stimulus has to reach a certain minimum level, specific to the particular organ, in order to evoke a geotropic bending response. The quantity of stimulus is equal to the gravitational force multiplied by the time for which it acts. For a given force, the period of exposure that is required to elicit a just detectable response is called the *presentation time*. Such threshold phenomena suggest that geoperception in plants involves the movement of free-falling bodies, or *statoliths*, which must move a certain distance to trigger the geotropic response mechanism. At a given temperature the presentation time is proportional to the inverse of the quantity of gravitational stimulus applied; that is, reciprocity holds for the geotropic stimulus as well as for the phototropic stimulus.

Mathematical analyses have been made to evaluate putative statoliths, and these have indicated that cellular inclusions as small as mitochondria could move rapidly enough within the cytoplasm in response to gravity to account for known presentation times. However, the kinetics of geoperception match most nearly the kinetics of gravity-induced displacement of starch grains in plant cells (Fig. 7.7). In fact, light- and electron-microscope studies of *statocytes* (gravity-sensing cells that contain statoliths) have revealed sedimentation of starch grains in response to gravity (Fig. 7.8), and also that the sedimentable starch grains are membrane-bound as *amyloplasts* (an amyloplast is a modified plastid, containing two to several starch grains). Not only are the positions of amyloplasts changed following re-orientation of statocyte cells, but other cytological events have been observed. Golgi bodies, for example, have been observed by Shen-Miller to sediment in geostimulated cells, and the distribution of the endoplasmic reticulum is also altered. However, although Golgi bodies and other organelles (in the *Chara* rhizoid barium sulphate crystals appear to serve as statoliths) may function as statoliths in some tissues, particularly those that do not contain amyloplasts, there is considerable circumstantial evidence that in most plant organs geotropic responses are initiated by the sedimentation of amyloplasts under the influence of gravity. Thus, plant organs deficient in starch (either naturally so or experimentally "destarched" by treatment with solutions of cytokinin and gibberellin, or low temperature) have been found to require very much longer presentation times for a geotropic response to occur.

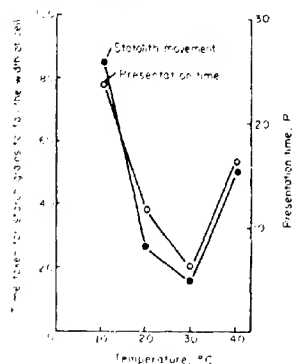


FIG. 7.7. Positive correlation observed between presentation time in geotropism of *Lathyrus odoratus* seedling stem, and time taken for starch grain sedimentation at different temperatures. (From L. Hawker, *Ann. Bot.* 47, 505-15, 1933.)

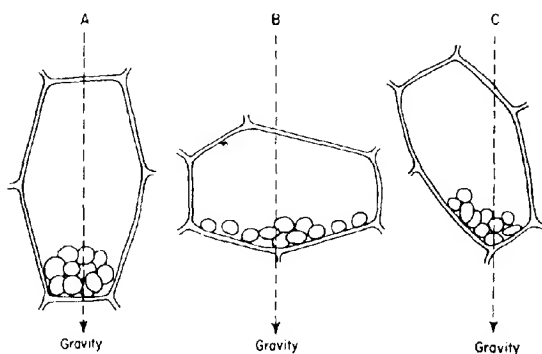


FIG. 7.8. Diagrammatic sections through statocytes following geotropic stimulation in three different positions. In each case the statoliths (amyloplasts) sedimented to the lowermost face of the cell. (From L. Hawker, *Ann. Bot.* 46, 121, 1932.)

Evidence that apical regions are the principal sites of geoperception in geotropic phenomena has been mainly derived from studies of grass and cereal coleoptiles, and of seedling roots of various monocotyledonous and dicotyledonous species. Removal of the apical few millimetres of either a coleoptile or a primary root prevents them from responding to a gravitational stimulus. Furthermore, replacing the tip with a layer of gelatine between stump and tip allows a geotropic curvature to develop, indicating that, as in phototropic

phenomena, perception of the stimulus takes place in the tip, following which there is evidently transmission of a chemical growth stimulus from the tip to the region of differential growth response.

In the case of geotropism in seedling roots, it has been established for several species that the important georesponsive amyloplasts are located in the central cylinder of root-cap cells. By micromanipulation it is possible to remove the root cap from otherwise intact roots, and this abolishes geotropic responsiveness in most species. Only roots that contain additional sedimentable amyloplasts in the root apical zone, or which rapidly form such amyloplasts after root-cap removal, retain some ability to respond to a gravitational stimulus after depriving them of the root cap.

Although it is quite well established that geoperception in geotropism involves sedimentation of statoliths, usually amyloplasts, in specifically situated statocyte tissues, little is known of the nature of the biochemical or physiological changes induced within statocyte cells by organelle displacement. Changes nevertheless certainly do take place, for after gravistimulation statocyte tissues affect the distribution of growth in the spatially separate responding region. In other words, the perceived stimulus results in transmission of some signal to the region of response.

The Mechanism of Redistribution of Growth in Geotropism

The tip regions of coleoptiles, stems and roots are not only sites of geoperception but also of the synthesis or release of growth hormones. Very little can be said about the means by which growth is redistributed in stems, largely because rather few studies have been made of geotropism in leafy shoots. Etiolated coleoptiles, in contrast, have received much more attention, and so too have seedling roots. Because the roles of growth hormones in roots are so little understood, it is necessary here to discuss separately available information on the geotropic response mechanisms in coleoptiles and roots. However, for both coleoptiles and roots it is generally considered that sedimentation of statoliths in statocyte cells results in compression of various cellular membranes, which in some unknown manner results in alterations in growth hormone synthesis, release and/or transport patterns.

Coleoptiles and stems. By analogy with the mechanism of response in phototropism, it might reasonably be expected that the differential growth occurring on the upper and lower sides of a horizontal organ involves an unequal distribution of auxin in the region of response, and there is much evidence in support of this conclusion. Various experiments have shown that auxin moves in the direction of the gravitational field, that is downwards. This means that when a plant is placed horizontally, higher auxin concentrations are created along the lower sides of the stem and root.

The first experiment demonstrating that auxin moves laterally downwards across a horizontal organ was performed in Holland by Dolk, in 1930, although the original proposal that this took place was put forward independently by Cholodny in 1924, and by

Went in 1926, Dolk placed excised coleoptile tips of *Avena sativa* and *Zea mays* horizontally, and collected the auxin which diffused from the upper and lower halves. More auxin was obtained from lower halves than from upper halves of tips of both species, while the total auxin yield of the organ was the same as that of similar but vertical tips. A number of other workers have repeated Dolk's experiments using various organs from other plant species, and found similar distributions of auxin between the top and bottom halves (Fig. 7.9). More recently it has been found that when radioactive IAA (IAA- ^{14}C) was applied to the apical ends of horizontal sections of the coleoptiles of *Zea mays* and *Avena sativa*, and of the hypocotyls of *Helianthus annuus*, it emerged asymmetrically into agar receiver blocks at the basal ends, so that more IAA- ^{14}C was found to have passed into the lower than into the upper receiver block.

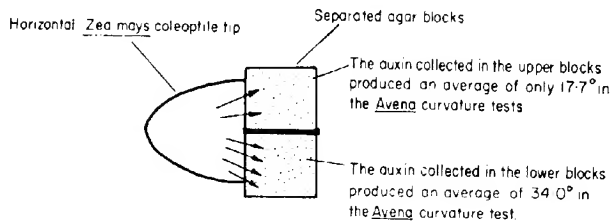


FIG. 7.9. Greater amounts of auxin diffuse from the lower side of a horizontally positioned coleoptile tip than from the upper side. The amount of auxin present in each agar block was measured by means of the Went *Avena* curvature test (see Chapter 3). (From B. Gillespie and W. R. Briggs, *Plant Physiol.* **36**, 364-8, 1961.)

Thus it has been clearly established that a lateral migration of auxin can occur under the influence of a gravitational field as well as of a light gradient. The higher concentration of auxin along the lower side of a hypocotyl or coleoptile presumably induces a greater growth in that region compared with the upper side, bringing about upward curvature.

Regardless of the nature of the geoperceptive system itself, gravity-stimulated tissues must possess a mechanism which brings about the lateral migration of auxin molecules. It is possible that this mechanism is an electrical one, for it has been found that a *geoelectrical potential* is set up across coleoptiles, stems and roots when they are positioned horizontally. Measurements of this electrical potential difference have shown that it amounts to from 5 to 20 millivolts, with the lower side positive with respect to the upper side. Thus, it is possible that the negatively charged ions of dissociated auxin move to the lower side under the influence of this electric field. However, recent work has shown that a lateral potential does not develop until curvature of the organ has commenced. This suggests that lateral displacement of auxin occurs *before* the geoelectrical potential is set up. Thus, it must be admitted that at present we have no idea how a displacement of starch grains in cells in the tip region could bring about a differential auxin distribution on the two sides of a horizontal organ.

Although there is well-substantiated evidence for the occurrence of downward lateral auxin transport in geostimulated coleoptiles, and limited amounts of similar data for hypocotyls, there is little or no convincing evidence for similar lateral transport of auxin in internodes. Very few experiments have been conducted on stem tissues other than hypocotyls, but these have tended to cast doubt upon the Cholodny–Went theory in that either no, or only very slight, downward lateral movement of radiolabelled auxin is detectable in horizontally positioned internode segments. It is, of course, possible that endogenous auxin levels rise in the lower sides of horizontal internodes of intact plants, but more careful research is needed to establish whether this is so.

Recently, endogenous gibberellins have been found to be asymmetrically distributed in geotropically responding internodes of sunflower (*Helianthus annuus*), with higher levels present in the lower, more rapidly elongating, side. Studies with radiolabelled gibberellins on stem segments has indicated that this asymmetry does not result from lateral displacement of gibberellin but possibly from greater synthesis and/or more rapid longitudinal transport in the tissues of the lower side of horizontal stems. Further work is required to evaluate the significance of these observations to an understanding of geotropism in stems.

Roots. Until recently, concepts of the response mechanism in root geotropism were similar in principle to the mechanism suggested by Cholodny and Went in the 1920s for phototropism and geotropism in coleoptiles. That is, it has been considered that auxin is synthesized or released from the root apex and undergoes lateral displacement towards the lower side of the organ. Because root elongation is inhibited by much lower concentrations of auxin than are supra-optimal for coleoptile or stem elongation (Chapter 5), it was thought that the auxin concentration on the lower side of a horizontally positioned root increases to become supra-optimal and consequently inhibitory to the growth of the auxin-sensitive root tissues. On the basis of this theory, the upper side of a horizontal orthogeotropic root would contain auxin at a more nearly optimal level and consequently grow at a more rapid rate than the inhibited lower side, resulting in a downward curvature of the root. However, current confusion as to the possible role of auxin in root elongation growth, and the problem of whether or not auxin is synthesized in root tips (Chapter 5), means that it is not possible to ascribe a role to auxin in root geotropism. Nevertheless, an open mind should be kept on the problem, for there exists experimental evidence that downward lateral displacement of auxin can occur in geotropically responding roots.

Over the past few years, it has been found that the root cap is not only the site of geoperception in root geotropic behaviour, but that in addition it appears to be a source of growth inhibitors, including abscisic acid, that play regulatory roles in root elongation. Both indirect and direct experimental evidence has been obtained to establish this. Thus, for example, M. B. Wilkins and his colleagues have done a range of experiments with *Zea mays* roots involving root-cap removal, removing half root caps, and the insertion of glass barriers (Fig. 7.10), which have convincingly demonstrated that the root cap exerts an inhibitory effect on root elongation and that under the influence of gravity this influence becomes asymmetrically distributed to cause greater inhibition of the lower half of a

horizontal root. Other direct analytical work has revealed the presence of several growth inhibitors in root caps, the most important of which appears to be abscisic acid (ABA). Preliminary studies of the movement of [^{14}C]-ABA have shown that ABA is transported basipetally in roots, and may be displaced laterally to the lower side in horizontal roots, but further more refined work is required to confirm this. On the basis of present evidence, therefore, it is reasonable to speculate that ABA may be concerned in the regulation of cell extension during the geotropic response of roots.

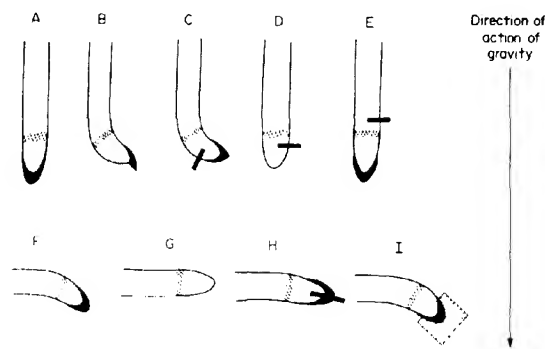


FIG. 7.10. Diagrammatic representation of some of the experiments which have indicated that the root cap is the source of growth inhibitor which is involved in the geotropic response mechanism in roots. The root cap is shown in black, and the elongation zone of the root is shaded. A. Vertical intact root grows downwards. B. Removal of half the root cap results in bending towards the remaining half cap regardless of the direction of gravity. C. Insertion of a glass barrier between half the root cap and the elongation zone has the same effect as removing half the cap. D. A similar barrier in the absence of the cap has no effect. E. A barrier positioned behind the growing zone is without effect. F. Intact horizontal root executing normal downward geotropic curvature. G. Removal of the root cap abolishes geotropism (because it appears to be both the region of geoperception and the source of growth regulating substances). H. A horizontal glass barrier through the root cap and apex abolishes, or largely removes, geotropism in a horizontal root. I. A glass barrier similar to that in H, but orientated vertically, does not prevent the development of a geotropic curvature. (Adapted from M. B. Wilkins, *Current Adv. Plant Sci.* 6(3), 317-28, 1975.)

The root cap thus possesses the capacity to detect gravity and also to produce and transport growth regulators in such a way as to control the direction in which the root grows. Furthermore, it appears that the root cap can also be sensitive to light, for in cereal roots the production of growth inhibitors, including ABA, by the cap rises following its exposure to light. This phenomenon may explain the fact that positive geotropism occurs in cereal roots only after they have been exposed to light, particularly as it has recently been found that addition of exogenous ABA to intact completely dark-grown roots of *Zea mays* induces downward bending in response to gravity.

Other Gravity-regulated Developmental Responses

We have considered above the physiology of orthogeotropism, especially as it is understood for coleoptiles and roots. However, for most plant organ geotropic behaviour is not necessarily constant. A given organ can change during development from being negatively to positively geotropic, or vice versa. Thus, in certain species flower and fruit stalks show such reversals between the flower-bud and mature fruit stages (e.g. in *Papaver*, *Fritillaria* and *Tussilago*). Correlative influences from other parts of the plant can also modify an organ's geotropic behaviour. The clearest example of this is seen in the influence that the apical bud of the main, orthogeotropic, shoot has upon the orientation of lateral organs such as leaves and lateral shoots. Lateral shoots and leaves are normally orientated at some angle between vertical and horizontal (i.e. they are plagiogeotropic), and removal of the apical bud of the main axis results in an upward (hyponastic) movement of both leaves and branches. One or more of the lateral branches usually become orthogeotropic and grow vertically upwards. Thus, it is clear that plagiogeotropic behaviour of laterals is at least partially determined by some correlative influence from the main apex. Exogenous auxin can substitute for the apical bud in maintenance of plagiogeotropism in laterals, which suggests that the correlative mechanism in regulation of geotropic behaviour is in some ways similar to that which operates in other apical dominance phenomena (Chapter 5).

Other important examples of the effects of gravity on plant development fall under the general heading of *gravimorphic effects*. The term *gravimorphism* has been used to categorize the morphogenetic, or developmental, effects that gravity can have in addition to geotropisms. Reaction-wood formation in plagiogeotropic branches is an obvious example of a gravimorphic effect. Others include the marked tendency for lateral buds to grow only from the upper sides of horizontal or plagiogeotropic shoots, the buds on the lower side remaining inhibited, and the promotion of flower-bud initiation in horizontally trained branches of fruit and other trees.

The physiology of these other types of gravity-induced growth and differentiation responses has received only limited study, and it is not yet possible to present a clear picture of mechanisms concerned. Nevertheless, from work that has been done, it appears that gravimorphism and plagiogeotropism are similar to orthogeotropism insofar as they are mediated through asymmetries in endogenous growth hormone distribution.

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CHAPTER 8

Phytochrome and Photomorphogenesis

IN THE preceding chapter we dealt with plant responses to the directional stimuli of light and gravity. In addition to phototropism, plants show other types of response to light signals. In the following chapter we shall consider plant responses to seasonal variations in the length-of-day (*photoperiodism*). In this latter type of response, the plant appears to have time-measuring capabilities which enable it to detect seasonal changes in the lengths of day and night. In addition to phototropism and photoperiodism, there are still other types of plant response to light signals, which are neither directional nor periodic, and which are included under the general term *photomorphogenesis*. As we shall see, photomorphogenesis includes a range of diverse phenomena which are controlled by specific photoreceptors, forming the *phytochrome* system.

THE RED/FAR-RED PHENOMENON

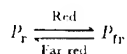
A major advance in our understanding of photomorphogenesis arose from the investigations of H. A. Borthwick and S. B. Hendricks of the U.S. Department of Agriculture, on the responses of a light-sensitive variety of lettuce seed. This type of seed shows little germination in the dark at 25° C, but germinates well if exposed to a short period of illumination. By exposing seeds to various parts of the spectrum Borthwick and Hendricks were able to demonstrate that the most effective region for the promotion of germination was in the red (maximum effectiveness at 660 nm), with a subsidiary peak in the blue. Far-red (maximum at 730 nm) radiation does not promote, but earlier work had shown that it is inhibitory to germination (Fig. 11.4). Borthwick and Hendricks investigated the interaction between the effects of red (R) and far-red (FR) radiation by exposing the seeds to R and FR alternately. They made the crucial discovery that the effects of R and FR are mutually reversible and that whether germination occurs or not depends on the nature of the last radiation to which the seeds are exposed (Table 8.1). Thus, each succeeding irradiation reverses the effect of the preceding treatment.

Now, it is clear that where a particular region of the spectrum causes a specific biological

TABLE 8.1. Control of lettuce seed germination by red and far-red light (from Borthwick *et al.*, *Proc. Nat. Acad. Sci. U.S.*, **38**, 662, 1952)

Irradiation	Percentage germination
Red	70
Red/Far-red	6
Red/Far-red/Red	74
Red/Far-red/Red/Far-red	6
Red/Far-red/Red/Far-red/Red	76
Red/Far-red/Red/Far-red/Red/Far-red	7
Red/Far-red/Red/Far-red/Red/Far-red/Red	81
Red/Far-red/Red/Far-red/Red/Far-red/Red/Far-red	7

effect, the tissues of the organism must contain a photoreceptor (or "pigment") which absorbs selectively in that region. Thus, it would appear that we have to postulate the presence of two photoreceptors in lettuce-seed tissue, one of which absorbs selectively in the red region and a second which absorbs in the far-red. However, Borthwick and Hendricks made the bold suggestion that there is essentially only one photoreceptor which can exist in the two alternative forms P_r and P_{fr} and that each form is capable of being reversibly converted into the other form, an hypothesis symbolized in the equation:



It will be seen that this hypothesis was based upon simple experiments on the effects of light upon the germination of lettuce seed. For some years the scheme remained entirely hypothetical, but later members of the same group constructed a special dual wavelength spectrophotometer which was capable of detecting small changes in the absorption spectra of etiolated plant tissues, at 660 nm and 730 nm. It was necessary to use etiolated plant tissues, since the presence of chlorophyll masks the absorption by other pigments in the red region. The instrument measured the *difference* in absorption of the tissues at 660 nm and 730 nm, during rapid alternation between irradiation at these two wavelengths, and it was found that after exposure of the dark-grown tissues to red light the absorption changed slightly, so that they absorbed more at 730 nm (Fig. 8.1), and the reverse change occurred if the tissues were now exposed to FR. Thus, the changes predicted by the hypothesis were fulfilled. In further work it was possible to demonstrate similar changes in cell-free extracts of etiolated tissues. Indeed it was possible to see a visible change in colour of the extracts with the naked eye, following exposure to R or FR. These changes in absorption properties were used to detect the presence of the photoreceptor(s) in further purification procedures and ultimately it proved possible to isolate a single protein which showed reversible changes in its absorption spectrum following exposure to R and FR, in exactly the manner originally postulated by Borthwick and Hendricks, who called this substance *Phytochrome*.

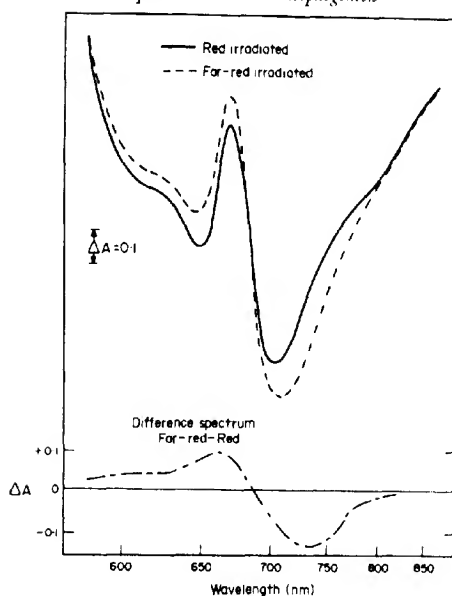


FIG. 8.1. "Difference spectrum" for phytochrome, showing absorbances of etiolated maize tissue after saturating exposures to red and far-red light (above) and the difference in absorbance (below). (From Butler, W. L., K. H. Norris, H. W. Siegelman and S. B. Hendricks, *Proc. Nat. Acad. Sci., U.S.A.* **45**, 1703, 1959.)

Phytochrome has subsequently been isolated in a very pure form and the protein part of the molecule apparently has a molecular weight of 120,000 daltons, while the non-protein, light-absorbing part (chromophore) has been shown to be a tetra-pyrrole compound related to the phycocyanins of the blue-green algae (Fig. 8.2). There is still some uncertainty as to the nature of the intra-molecular changes undergone by the chromophore when exposed to R or FR, but one suggestion is illustrated in Fig. 8.2. There is also some evidence that the protein part of the molecule may undergo a conformational change during photoconversion. The absorption spectra of pure P_r and P_{fr} are given in Fig. 8.3.

THE RANGE OF PHYTOCHROME-CONTROLLED RESPONSES

Since the original discovery of R/FR reversibility in lettuce seeds, similar effects have been demonstrated for a wide variety of plant responses and in all the main groups of plants from the green algae to the flowering plants (Table 8.2). It will be seen that these

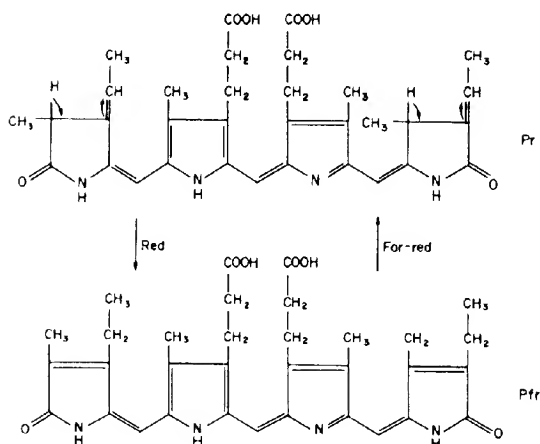


FIG. 8.2. Suggested changes within the chromophore of phytochrome occurring during photoconversion. (From H. W. Siegelman, D. J. Chapman and W. J. Cole, in *Porphyrins and Related Compounds*, Ed. T. W. Goodwin, Academic Press, London, 1968.)

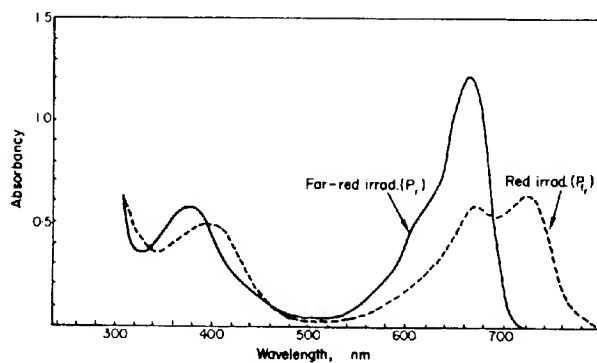


FIG. 8.3. Absorption spectra of a solution of oat phytochrome following irradiation with red and far-red light, giving the P_{fr} (broken line) and P_r (continuous line) forms, respectively. (From H. W. Siegelman and W. L. Butler, *Ann. Rev. Plant Physiol.* **16**, 383, 1965.)

TABLE 8.2. Some phytochrome-controlled responses

<i>Algae, bryophytes and pteridophytes</i>	<i>Angiosperms</i>
Spore germination	Seed germination
Chloroplast movement	Hypocotyl hook formation
Protonema growth and differentiation	Internode extension
	Root primordia initiation
	Leaf initiation and growth
	Leaflet movement
	Electrical potential
	Membrane permeability
	Phototropic sensitivity
	Geotropic sensitivity
	Anthocyanin synthesis
<i>Gymnosperms</i>	
Seed germination	
Hypocotyl hook formation	
Internode extension	
Bud dormancy	

responses include spore germination, epicotyl hook opening, leaf expansion, internode extension, and root initiation, as well as numerous responses at the sub-cellular and molecular level, such as chloroplast movement, enzyme synthesis and changes in membrane permeability. In all these cases it has been demonstrated that R and FR have opposite effects, with similar action spectra (Fig. 8.4), and show R/FR reversibility. These R/FR effects are so characteristic of phytochrome that where they can be demonstrated for any given biological response, the latter can be assumed to involve phytochrome. It is these manifold aspects of growth and development under phytochrome control which are referred to as *photomorphogenesis*.

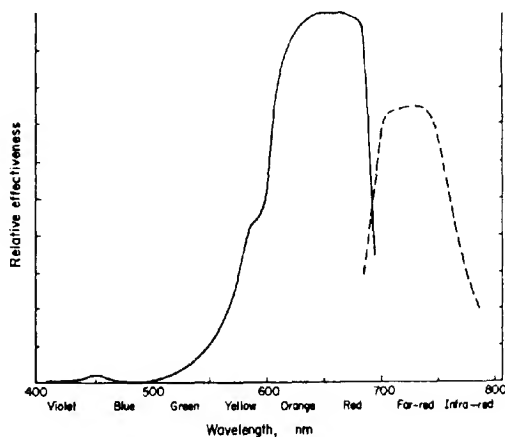


FIG. 8.4. Action spectra for the red and far-red physiological responses controlled by phytochrome. Continuous line: red effects; broken line: far-red effects. (Adapted from F. B. Salisbury, *Endeavour*, 24, 78-80, 1965.)

It is evident that phytochrome control does not apply only to special phenomena, such as light sensitivity in seeds, but is involved in some of the most general aspects of development, such as leaf expansion and stem extension in normal development of the green shoot. Everyone is familiar with the characteristic appearance of etiolated shoots which have grown in complete darkness. These symptoms of etiolation can be reduced by quite short periods (5 minutes) of daily irradiation with red light, and the effects of R can be reversed by FR, indicating phytochrome control (Fig. 8.5). However, the development of chlorophyll requires longer periods of irradiation and we shall see that the full development of what we regard as a "normal" green shoot requires quite high energy levels (p. 196).

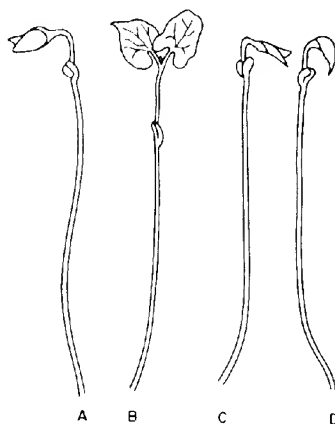


FIG. 8.5. Phytochrome control of shoot development in bean (*Phaseolus vulgaris*). Treatments: (a) grown in continuous darkness; (b) exposed to 2 minutes red light; (c) 2 minutes red and 5 minutes far-red; (d) 5 minutes far-red. (From R. J. Downs, *Plant Physiol.* **30**, 468, 1955.)

DETECTION AND MEASUREMENT OF PHYTOCHROME IN VIVO

The detection and measurement of phytochrome in plant tissues is based upon spectrophotometric measurements of the differences in light-absorption spectra at 660 nm and 730 nm following irradiation with R and FR, using similar methods to those employed for the original isolation of phytochrome. The tissue is first irradiated with red light to convert the phytochrome to the P_{fr} form and the difference between the absorbance at 660 nm and at 730 nm determined (ΔA_r). The pigment is then converted to the P_r form by exposure to FR, and the difference in absorbance at the two wavelengths again determined (ΔA_{fr}).

The overall difference between the two measurements (Fig. 8.6) will be related to the total amount of phytochrome present.

That is:

$$\Delta(\Delta A) = \Delta A_{tr} - \Delta A_r$$

where ΔA is the difference in absorbance at 660 and 730 nm.

Etiolated tissues have been found to have a high content of phytochrome, the highest concentrations being found in meristematic tissues, including shoot and root tips and cambial tissue. The phytochrome content of green tissue is too low to detect spectrophotometrically, but low amounts have been found in extracts of leaves of a number of species.

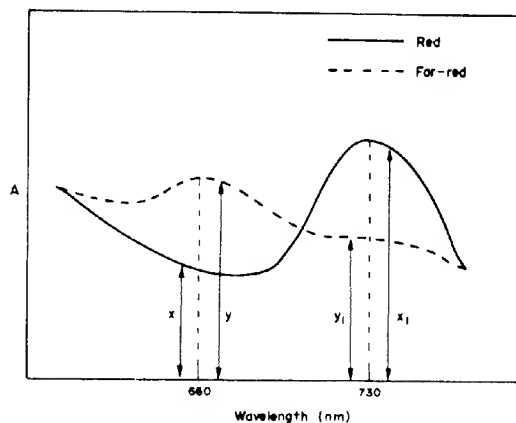


FIG. 8.6. Diagram of absorbance changes measured in the phytochrome assay. Note: $\Delta_r = x - x_1$ (after red), $\Delta_{tr} = y - y_1$ (after far-red). (From R. E. Kendrick and B. Frankland, *Phytochrome and Plant Growth*, Edward Arnold, London, 1976.)

THE INTRACELLULAR LOCALIZATION OF PHYTOCHROME

Attempts have been made to identify the sites at which phytochrome occurs within the cell. The most direct approach has been through the use of immunocytochemical techniques. Rabbit antibodies against purified phytochrome were produced and applied to sections of plant tissues. As a result, rabbit antibody molecules will become attached to phytochrome molecules within the cell. The section is then treated with sheep antiserum to rabbit antibody and this is followed by a rabbit antiperoxidase-peroxidase complex so that each phytochrome molecule is "tagged" with the enzyme peroxidase, the location of which can then be detected by histochemical methods (Fig. 8.7). Such studies have indicated that in the P_r form phytochrome is not strictly localized within the cell, and occurs in

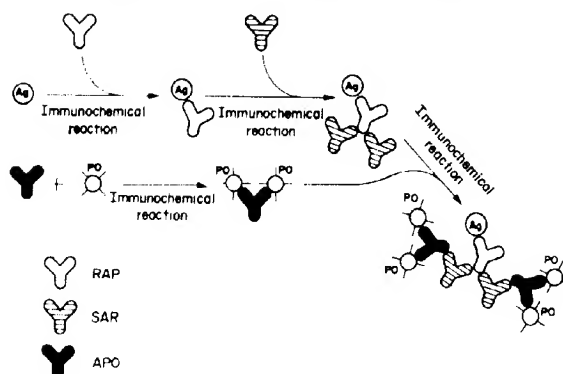


FIG. 8.7. A double indirect technique for the visualization of an antigen (see text). Components of the labelling procedure are: Ag, phytochrome; RAP, rabbit anti-phytochrome serum; SAR, sheep anti-rabbit immunoglobulin serum; APO, rabbit anti-peroxidase immunoglobulins; PO, peroxidase. (From L. H. Pratt, R. A. Coleman and J. M. Mackenzie, Jr., in *Light and Plant Development*, Ed. H. Smith, Butterworths, London, 1976.)

mitochondria and plastids, as well as in the cytoplasm generally, but not in nuclei or vacuoles. However, on conversion to the P_{tr} form by exposure to R light, phytochrome rapidly becomes localized in discrete areas which have not yet been identified.

Other evidence indicates that the phytochrome is located in various cell membranes. Thus, cell fractionation techniques including high-speed centrifugation, have yielded fractions consisting of cell membranes and containing a high proportion of the total cell phytochrome. The association of the phytochrome with the membrane occurs only in tissues pre-treated with red light, suggesting that it is only the P_{tr} form which is attached to the membranes. Other cell fractionation techniques have demonstrated that phytochrome is present in etioplasts of etiolated wheat and barley leaves.

A different approach to the problem of locating phytochrome within the cell has been used in studies on movements of the chloroplasts of the filamentous green alga, *Mougeotia*. Each cell of this alga contains a single, plate-like chloroplast which turns so that it is edge-wise to the direction of the incident light at high intensities but at right angles to the light at low intensities. By using microbeams of R and FR light, it has been shown that these chloroplast movements involve phytochrome, and that the response can be obtained when only the outer layers of the cytoplasm are irradiated. Moreover, using microbeams of polarized R and FR, it was shown that R is only effective when the plane of the electric vector is *parallel* to the cell surface and that FR is only effective when the plane of the electric vector is *at right angles* to the cell surface (Fig. 8.8). This finding indicates that the phytochrome molecules are arranged in a regular manner at or near the cell surface and that photoconversion involves a change in orientation of the molecules (Fig. 8.9). We shall

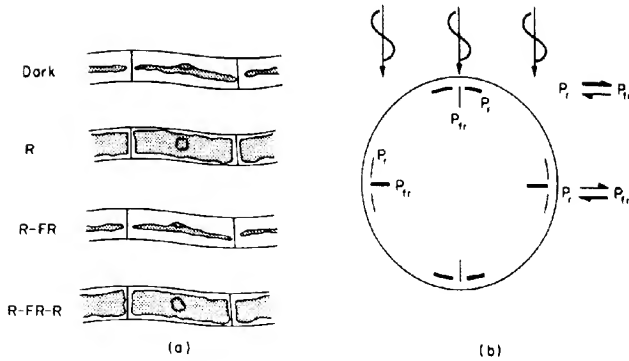


FIG. 8.8. (a) Induction of chloroplast orientation in *Mougeotia* by red light (R) and reversion of red effect by far-red (FR), starting from the profile orientation (above). (b) Absorption of polarized light by phytochrome molecules which change dichroic orientation with conversion $P_r \rightarrow P_{fr}$. Cross-section of a *Mougeotia* cell with P_r (surface parallel dashes) and P_{fr} (surface normal dashes). Molecules which are in favourable geometric position to absorb polarized light are heavy lined. As a consequence, the photostationary state is shifted, according to the position, to the left or right side (heavy-lined arrows). (Note that the electric vector is at right angles to the direction of the incident polarized light, as indicated by the sinusoidal curves above.) (From W. Haupt, in *Phytochrome*, Ed. K. Mitrakos and K. Shropshire, Academic Press, London, 1972.)

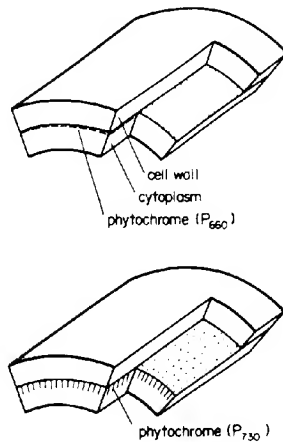


FIG. 8.9. Suggested model showing the orientation of phytochrome molecules in the plasma membrane (top) for the P_r form and (below) for the P_{fr} form. (From W. Haupt, *Phytochrome*, Ed. K. Mitrakos and K. Shropshire, Academic Press, London, 1972.)

see later that other evidence suggests that phytochrome acts by affecting the permeability of cells, again suggesting that it acts at the cell membrane.

PHOTOCONVERSION OF PHYTOCHROME IN THE CELL

We have seen that the P_r and P_{fr} forms of phytochrome can readily be interconverted by exposure to R and FR respectively. When the R/FR reversibility phenomena was first discovered it was believed that after conversion of P_r to P_{fr} by exposure to red light, reversion of P_{fr} to P_r could take place spontaneously in the dark, without exposure to FR. However, such "dark reversion" has only been demonstrated in certain dicotyledons (e.g., in cauliflower tissue (Fig. 8.10)) and has not yet been shown to occur in monocotyledons.

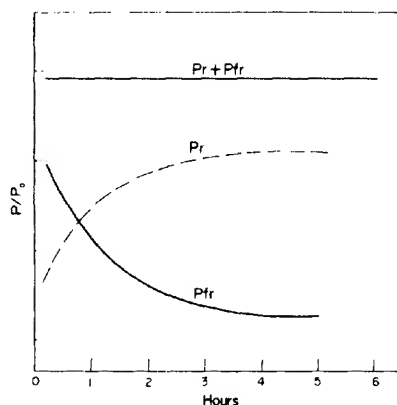


FIG. 8.10. Dark reversion of P_{fr} to P_r in the complete absence of P_{fr} destruction in cauliflower curd tissue after 5 minutes red-light. (From W. L. Butler, H. C. Love and H. W. Siegelman, *Plant Physiol.* **38**, 514, 1963.)

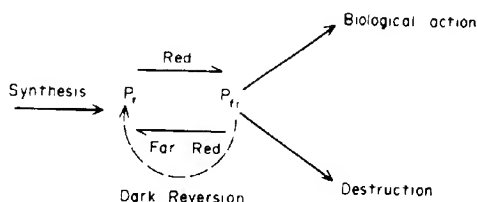
Nevertheless, the level of P_{fr} generally declines quite rapidly in most plant tissues and concomitantly the total phytochrome (P) in the tissues also declines. This decline in P_{fr} can be arrested by exposure to FR, i.e. by conversion to P_r . Thus, whereas P_r is relatively stable, P_{fr} is highly labile and is evidently destroyed quite rapidly *in vivo*. It is not understood how such destruction of P_{fr} occurs, but evidently P_{fr} is the biologically active form and it must be involved in the initiation of the chain of processes which are ultimately manifested in an observable biological response; it may be that the action of P_{fr} entails its destruction.

In most experiments involving R/FR reversibility, the FR treatment follows immediately after exposure to R, but if exposure to FR is delayed for increasing periods a point is

reached when FR no longer reverses the effect of R, indicating that the events initiated by conversion to P_{tr} have already been set in progress. This period beyond which reversibility is lost is referred to as the "escape" time and varies from 1.5 minutes to 9 hours in different tissues.

If plant tissues are allowed to remain in the dark following a single exposure to R, the initial decline in total phytochrome is later followed by an increase, suggesting that synthesis of new phytochrome in the P_t form takes place. This synthesis has been demonstrated by density-labelling (see p. 90) by demonstrating the incorporation of deuterium into P_t when plant tissues are incubated in D_2O .

As a result of the various processes described above, it is clear that the phytochrome in active plant tissues undergoes continuous "turnover", involving synthesis, interconversion, destruction and reversion, which may be symbolized as follows:



MODE OF ACTION OF PHYTOCHROME

Many phytochrome-controlled responses appear to involve the action of genes which previously were not expressed. For example, the development of epidermal hairs and of anthocyanin in mustard hypocotyls is under phytochrome control and would appear to involve the expression of genes which are not expressed in dark-grown seedlings. Moreover, gene expression involved both enzyme synthesis and enzyme activity, and phytochrome control of both these processes has been demonstrated (p. 194). These observations might seem to indicate that phytochrome acts by controlling either enzyme synthesis at the transcription or translation level, or by controlling the activation of pre-existing enzymes. However, any attempt to formulate a general hypothesis to account for the very varied range of phytochrome-controlled responses must take into account various effects which appear to involve rapid changes in membrane permeability.

For example, the leaves of certain plant species, such as *Mimosa pudica* and *Albizia julibrissin*, show "sleep movements" involving upward or downward folding of the leaflets. These movements, which involve changes in the turgor pressure in cells of the pulvini at the point of attachment of the leaflet to the midrib, are evidently under some degree of phytochrome control, since they are inhibited by exposure to FR at the end of the light period and this effect can be reversed by R. These effects of FR can be observed

within about 10 minutes. The changes in turgor have been shown to be accompanied by the movement of potassium ions into and out of the cell, thereby affecting its osmotic properties. An even more rapid response is seen in root tips of barley and mung bean, which, under certain conditions, will adhere to a negatively charged glass surface following exposure to R light and this effect can be reversed by FR. This electrostatic effect indicates that the root surface becomes positively charged on exposure to R light and that this effect is reversed by FR.

Exposure to red light has also been shown to result in rapid changes in electrical potential in several organs. For example, exposure of etiolated coleoptiles to R light causes the tip to become more electropositive with respect to the base within 15 seconds, and similar changes have been observed in mung bean root tips. It seems clear that these various types of rapid effect following phytochrome conversion involve changes in the permeability properties of the cell membranes to electrolytes such as potassium ions.

Since these rapid effects may occur almost instantaneously or within a few minutes, the time involved is too short to depend upon RNA or protein synthesis, for which a period of at least 15 minutes would appear to be required. Hence, it is generally held that the primary mode of action of phytochrome must involve changes in cell membrane properties, a conclusion which is consistent with the evidence of the intracellular localization of phytochrome (p. 190).

However, although the *primary* mode of action may involve changes in membrane permeability, there is good evidence that the secondary effects of phytochrome conversion involve enzyme-controlled processes. Thus, in *Avena* seedlings irradiation with R light results in a rapid conversion of ADP to ATP, and in other tissues there are changes in the levels of NADP which can be prevented by FR irradiation. Moreover, exposure to R light leads to increased activity of a wide range of enzymes. Thus, R light increases the level of phenylalanine ammonia lyase (PAL) (Fig. 8.11) which catalyses the removal of ammonia from phenylalanine to give cinnamic acid, in mustard seedlings. In this instance R appears to lead to activation of pre-existing enzyme rather than to new enzyme synthesis, but in other tissues R appears to stimulate the synthesis of ascorbic acid oxidase. Irradiation of etiolated bean leaves with R light leads to marked increases in polyribosomes within 1 hour, suggesting that the treatment results in increased availability of messenger RNA.

The nature of the processes connecting the primary changes in membrane permeability with the diverse metabolic events involving phytochrome remain obscure. However, there is now considerable evidence that exposure to R light results in rapid increases in extractable gibberellins and cytokinins, in etiolated and green leaves and in certain seeds. Thus, exposure to 5 minutes of R light causes marked increases in gibberellins in etiolated barley and wheat leaves (Fig. 8.12), and in cytokinin levels in *Rumex* seeds and leaves of poplar (*Populus robusta*). Thus, it is possible that these hormones act as "second messengers" and provide a link between the primary and secondary effects of phytochrome conversion. For example, etiolated barley and wheat leaves grown in complete darkness are tightly rolled, but they can be induced to unroll by exposure to a short period of red light, the effect of which can be nullified by far red. However, sections of etiolated barley leaves can

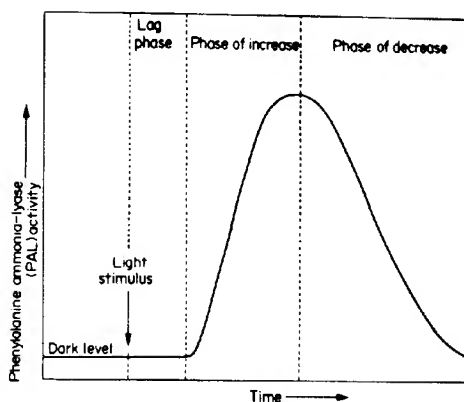


FIG. 8.11. Generalized *in vivo* response pattern of phenylalanine ammonia-lyase (PAL) activity to irradiation, showing the lag phase, phase of increase and phase of decline. (From H. Smith, *Phytochrome and Photomorphogenesis*, McGraw-Hill, London, 1975.)

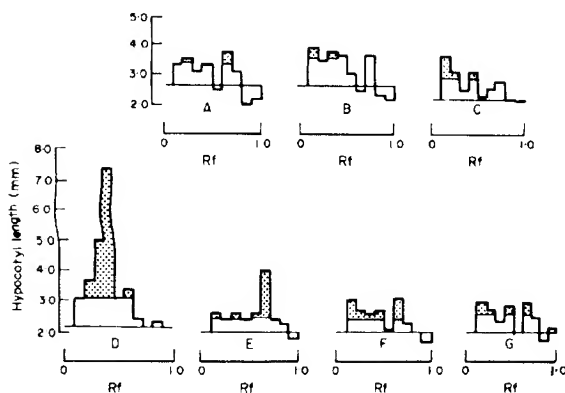


FIG. 8.12. Effect of red light on endogenous gibberellin levels in etiolated wheat leaves. Each separate histogram represents the gibberellin activity present in extracts of leaves after separation into ten fractions by paper chromatography. Each fraction was tested by its promoting effect on the growth of lettuce hypocotyls. The solid parts of the histograms represent gibberellin activity significantly above control

A = dark-grown leaves,

E = 5 minutes red light + 20 minutes dark,

B = 5 minutes red light,

F = 5 minutes red light + 30 minutes dark,

C = 5 minutes red light + 5 minutes dark,

G = 5 minutes red light + 60 minutes dark,

D = 5 minutes red light + 10 minutes dark,

(From L. Beevers, L. B. Loveys, J. A. Pearson and P. F. Wareing, *Planta (Berl.)*, **90**, 286, 1970).

be induced to unroll in darkness by application of exogenous GA_3 suggesting that the red-light effects may be mediated by the increases in endogenous gibberellins.

THE "HIGH IRRADIANCE REACTION"

As we have seen, the characteristic features of etiolated shoots grown in complete darkness (viz. excessive internode elongation, poor leaf development, lack of chlorophyll, etc.) can be greatly reduced by exposing them to quite short periods (e.g. 5 minutes) of R light each day, and the effect of R can be reversed by FR, clearly indicating involvement of phytochrome. However, the appearance of shoots exposed to only short periods of low-intensity R light in this way is still far from "normal" and in order to obtain the typical appearance of shoots grown in natural daylight it is necessary to expose the shoots to several hours of daylight at high intensity each day. Now, work on white mustard seedlings showed that whereas short periods of FR at low intensity reversed the effect of R, longer periods of FR produced the *same* effects as R, i.e. they reduce the characteristic symptoms of etiolation by causing expansion of the cotyledons, reduction of hypocotyl extension and the development of anthocyanin. Moreover, whereas the effects of short periods of FR reached "saturation" at quite low intensities, the effects of longer periods of FR continued to increase as the intensity was increased to high levels. Hence these effects were said to involve a "High Irradiance Reaction" (HIR).

Action spectra for the HIR show major peaks in the FR at 710-730 nm, and in the blue region (Fig. 8.13). At first it was thought that the HIR must involve a separate photoreceptor, other than phytochrome. However, later studies have indicated that HIR effects in the

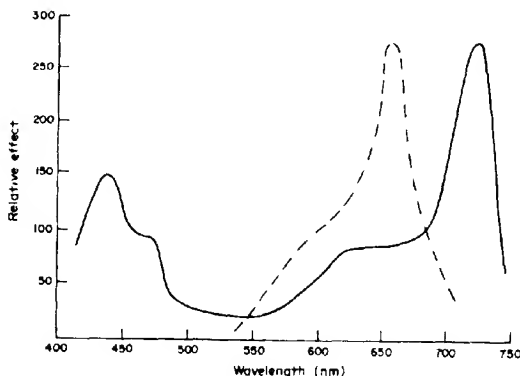


FIG. 8.13. Generalized action spectrum for the high-irradiance reaction (solid line), together with that of the low-irradiance phytochrome reaction (broken line). (From H. Mohr, *Biol. Rev.* 39, 87, 1974.)

FR region are almost certainly mediated via phytochrome. The evidence for this conclusion is based on the observation that when plant tissues are exposed to FR, 3 per cent of the total phytochrome remains in the P_{fr} form. This is because the absorption spectra of P_r and P_{fr} overlap (Fig. 8.3), so that when tissue is exposed to FR, a "photostationary state" is reached, representing an equilibrium between photoconversion of P_{fr} to P_r and the reverse reaction. It was shown that the effects of the HIR could be simulated by irradiating tissues with mixtures of R and FR at various intensities and in varying proportions, such that 3 per cent of the total phytochrome was present in the P_{fr} form. Thus, it appears that phytochrome is the photoreceptor for the high irradiance FR effects. However, there is evidence that the effects in the blue region of the action spectrum for the HIR involve a different photoreceptor. Thus, the action spectrum in the blue region of the HIR is identical with that for phototropism in both higher plants and in fungi, which do not contain phytochrome. Again, gherkin seedlings transferred from dark to blue light show a reduced rate of hypocotyl extension after a lag of 30 seconds, whereas with far red the lag is 40 minutes. The photoreceptor involved in the blue region of the HIR is probably a flavin, responsible for many photoresponses in both lower and higher plants.

Under natural conditions plants are not, of course, subjected to monochromatic R and FR but normal daylight will induce a "photostationary state" in which a significant proportion of the total phytochrome will be in the P_{fr} form. From the evidence presented above it would appear that the HIR of phytochrome is involved in normal development of green shoots, under natural conditions.

Another aspect of the role of phytochrome under natural conditions arises where plants occur at high density, both in natural vegetation and in crops, so that they naturally shade each other. Natural daylight contains approximately equal intensities of R and FR, but because chlorophyll absorbs strongly in the red region, the light within a leaf canopy will contain a much higher proportion of FR (Fig. 8.14). It has been shown that radiation

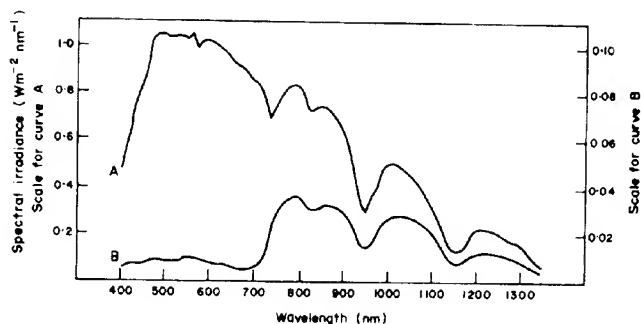


FIG. 8.14. Spectral distribution of light in the open at noon on a sunny day in July (A), compared with that of light within a woodland (B). (From R. E. Kendrew and B. Frankland, *Phytochrome and Plant Growth*, Edward Arnold, London, 1976.)

containing a high proportion of FR tends to lead to increased internode elongation and this may well explain the abnormal stem elongation frequently observed in dense stands of plants, and which must confer an advantage on plants with this capacity, in the intense competition for light. The role of phytochrome in the dormancy of light-sensitive seeds and in their responses under natural conditions are discussed in Chapter 11.

FURTHER READING

General

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More Advanced Reading

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CHAPTER 9

The Physiology of Flowering—

I. Photoperiodism

INTRODUCTION

In the "typical" life history of a herbaceous flowering plant, there is usually an initial phase of vegetative growth, which sooner or later is followed by the reproductive phase. However, there is a good deal of variation with respect to the distinctness of these two phases; in some plants there is a fairly sharp transition from vegetative growth to reproduction, as in wheat or sunflower, while in others vegetative growth and flowering occur concurrently, as in runner beans (*Phaseolus multiflorus*) or tomato (*Lycopersicon esculentum*). In the first type of plant, growth is usually *determinate*, the axis being terminated by an inflorescence, whereas in the second group the flowers are axillary and borne on lateral shoots, while the main axis continues vegetative growth. In plants of both groups, however, there is nearly always a certain minimum period of purely vegetative growth—only very exceptionally can flowers be formed immediately following germination, e.g. in *Chenopodium rubrum*, under shortday conditions. The duration of this vegetative phase is very variable. Usually there is a certain period of growth during which a succession of new leaves is formed at the shoot apex. In some perennial species, however, the number of leaves is already predetermined in the stage of dormancy and the vegetative phase involves only the expansion of leaf primordia already laid down in the previous year, as in many bulbous plants and woody species.

Factors Determining the Onset of Flowering

What causes the transition from the vegetative to the reproductive phase? Is it due to some internal control mechanism, determined by the genetical make-up of the species, or is it dependent upon a change in external conditions? Now, plants differ very greatly in their sensitivity to external conditions, the development of some species being relatively insensitive, so that provided that the environmental conditions are not so unfavourable that

growth is completely prevented, they will ultimately flower under a wide range of conditions. In other species, however, the initiation of flowering is very sensitive to external conditions and will not occur under certain conditions, e.g. of temperature or daylength, even though these may be quite favourable for growth; that is to say, the requirements for flowering are not necessarily the same as for vegetative growth.

Since our knowledge of the physiology of flowering is much more complete for species which are sensitive to environmental conditions, we shall first deal with this group, and then describe briefly what is known regarding the control of flowering in the "insensitive" group.

Quantitative Measurement of Flowering Responses

The difference between the vegetative and the flowering condition is a qualitative one. It is important, however, that in studying the physiology of flowering, we should have an exact measure of the flowering responses to various treatments. Various indices of flowering have been used, such as (1) the percentage of flowering plants in the total group receiving a particular treatment; (2) the total number of flowers or total number of flowering nodes; (3) the time to the first appearance of flowers (the shorter the time the greater the flowering response); (4) the number of leaves formed before flower initiation; (5) the use of a scale of "scores" depending upon the stage of development reached by the flowers. This latter method is used where the flowers formed are still microscopic and incompletely developed. An arbitrary scale of "scores" is assigned to various stages of development (Fig. 9.1) and the total "score" of a given batch of plants is determined.

PHOTOPERIODISM

The fact that seasonal changes in daylength conditions profoundly affect the life cycle of many plants was first clearly demonstrated by Garner and Allard, two American plant breeders, in 1920. Garner and Allard were originally concerned with the peculiar seasonal flowering behaviour of certain varieties of tobacco (*Nicotiana tabacum*) and soybeans (*Glycine max*). A newly developed variety of tobacco, Maryland Mammoth, was found to grow vigorously throughout the summer, but did not flower. When grown in a greenhouse during the winter, this same variety flowered and fruited abundantly. With certain varieties of soybeans, plantings at successive intervals during the spring and summer all tended to flower at the same date in the late summer, the vegetative period being progressively shortened the later the date of sowing. Garner and Allard attempted to regulate the flowering of the tobacco and soybeans by varying the temperature, nutrition and soil moisture, but none of these factors was found to affect the date of flowering very markedly. They then investigated the effect of shortening the daily light period by a few hours by placing the plants in a dark chamber, and found that under the shortened daylight period

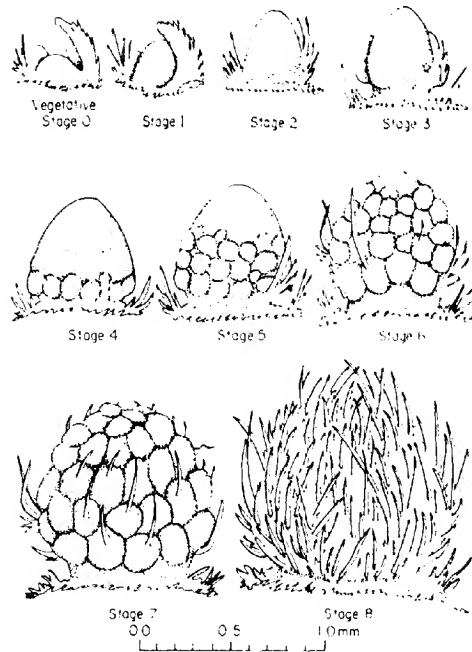


FIG. 9.1. Stages of development of the staminate inflorescence primordium of *Xanthum*.
(From F. B. Salisbury, *Plant Physiol.* **30**, 327, 1955.)

the plants quickly initiated flowers. After this exciting discovery, they proceeded to test the effect of various daylength conditions on a wide variety of plant species. Variation of daylength was achieved in two ways: (1) during the summer months by shortening the natural daylength, and (2) in winter by extending the natural daylength by artificial illumination.

It was found that for many plant species the daylength (i.e. the lengths of daily light and dark periods) is a very important factor in growth and development, particularly in the control of flowering, and the phenomenon is known as *photoperiodism*. On the other hand, in other species, flowering was not materially affected by length of day. The group in which daylength has a marked effect may be separated into two subdivisions, viz. (1) those in which flowering is readily induced by exposure to short days, and known as "*short-day*" plants (SDP), and (2) a second group in which flowering is favoured by long days, and which are known as "*long-day*" plants (LDP). Species in which daylength does not markedly affect flowering are known as *indeterminate* or "*day-neutral*" plants.

Some species remain permanently vegetative if kept under unfavourable daylength

conditions, and hence may be called obligate photoperiodic plants. They include both SDP, such as *Xanthium pennsylvanicum*, and LDP such as *Hyoscyamus niger*. Other species may show hastened flowering under SD or LD, but they will ultimately flower even under unfavourable daylength conditions, and hence show a *quantitative* photoperiodic response. Such species include the SDP *Salvia splendens*, rice (*Oryza sativa*) and cotton (*Gossypium hirsutum*) and the LDP wheat (*Triticum*) and flax (*Linum usitatissimum*). Obligate photoperiodic plants show a well-marked *critical daylength*, below or above which flowering will not occur in LDP and SDP, respectively, whereas facultative photoperiodic species show only a graded response to daylength, with no sharp cut-off point. Examples of SDP and LDP are shown in Fig. 9.2, and are listed in Table 9.1.

TABLE 9.1. Some examples of short-day and long-day plants

Short-day plants
A. Species with an Absolute or Qualitative Short-day Requirement

<i>Amaranthus caudatus</i> (Love-hes-bleeding)	<i>Ipomoea hederacea</i> (Morning glory)
<i>Chenopodium album</i> (Pigweed)	<i>Kalanchoe blossfeldiana</i>
<i>Claytonia perfoliata</i> (Poinsettia)	<i>Lemna perpusilla</i> (Duck weed)
<i>Coffea arabica</i> (Coffee)	<i>Nicotiana tabacum</i> (Tobacco, var. Maryland Mammoth)
<i>Euphorbia pulcherrima</i> (Poinsettia)	<i>Perilla ocymoides</i>
<i>Fragaria</i> (Strawberry)	<i>Xanthium strumarium</i> (Cocklebur)
<i>Glycine max</i> (Soybean)	

B. Species with Quantitative Short-day Requirement

<i>Cannabis sativa</i> (Hemp)	<i>Oryza sativa</i> (Rice)
<i>Cosmos bipinnatus</i> (Cosmos)	<i>Saccharum officinarum</i> (Sugar cane)
<i>Gossypium hirsutum</i> (Cotton)	<i>Salvia splendens</i>

Long-day plants**A. Species with an Absolute or Qualitative Long-day Requirement**

<i>Alopecurus pratensis</i> (Foxtail grass)	<i>Melilotus alba</i> (Sweet clover)
<i>Anagallis arvensis</i> (Pimpernel)	<i>Mentha piperita</i> (Peppermint)
<i>Anethum graveolens</i> (Dill)	<i>Phleum pratensis</i> (Timothy grass)
<i>Avena sativa</i> (Oat)	<i>Raphanus sativus</i> (Radish)
<i>Dianthus superbus</i> (Carnation)	<i>Rudbeckia bicolor</i> (Coneflower)
<i>Festuca elatior</i> (Fescue grass)	<i>Sedum spectabile</i> (Sedum)
<i>Hyoscyamus niger</i> (Henbane)	<i>Spinacia oleracea</i> (Spinach)
<i>Lolium temulentum</i> (Rye-grass)	<i>Trifolium</i> spp. (Clover)

B. Species with a Quantitative Long-day Requirement

<i>Antirrhinum majus</i> (Snapdragon)	<i>Petunia hybrida</i> (Petunia)
<i>Beta vulgaris</i> (Garden beet)	<i>Pisum sativum</i> (Garden pea)
<i>Brassica rapa</i> (Turnip)	<i>Poa pratensis</i> (Kentucky blue-grass)
<i>Hordeum vulgare</i> (Spring barley)	<i>Secale cereale</i> (Spring rye)
<i>Lactuca sativa</i> (Lettuce)	<i>Triticum aestivum</i> (Spring wheat)
<i>Oenothera</i> spp. (Evening primrose)	



FIG. 9.2. Plants of *Kalanchoë blossfeldiana* (left) and of *Rudbeckia bicolor* (right), flowering in response to short days and long days, respectively.

The short-day group includes many plants which are indigenous to regions of low latitude, north or south of the equator, such as rice, sugar cane, hemp, millet and maize, where the daylength never exceeds more than 14 hours at any season of the year. SDP of temperate regions, where the days are long in the summer, usually initiate flowers only in the late summer as the days shorten, e.g. the cultivated "Michaelmas daisies" (*Aster* spp.) and *Chrysanthemums*. The typical LDP are native to the temperate regions, and flower under the naturally long days of summer. They include many of the grasses and cereals, and other common cultivated plants such as spinach (*Spinacia oleracea*), lettuce (*Lactuca sativa*), beet (*Beta vulgaris*), flax (*Linum usitatissimum*) and clover (*Trifolium* spp.), as well as many wild species. In addition to the two main groups of SDP and LDP, there is a smaller number of species with dual daylength requirements. Thus, certain species require to be exposed first to LD and then to SD for flower initiation to occur and hence are called "long-short-day" plants (LSDP); examples of this type of response are provided by *Bryophyllum crenatum* and *Cestrum nocturnum*. Other species, such as *Scabiosa succisa*, *Campanula medium* and *Trifolium repens*, require to be exposed first to SD and then to LD, and hence are called "short-long-day" plants (SLDP).

Where a species has a wide distribution, so that there is a considerable difference in latitude between its northern and southern limits, it is found that it is differentiated into a number of races or ecotypes, differing in their daylength responses, e.g. golden rod (*Solidago sempervirens*), which has a wide distribution along the western coast of North America, and perennial ryegrass (*Lolium temulentum*), which has a wide distribution in Europe and North Africa. These different forms within a given species usually show a closely graded

series, from typical SDP at one end to LD-tolerant types at the other, or from typical LDP to SD-tolerant types.

Other Responses Affected by Daylength

In addition to the onset of flowering, daylength may affect certain other purely vegetative processes in the plant. Thus, it is frequently found that the length of the internode may be much reduced under SD as compared with LD. This effect is seen at its extreme form in certain long-day species, which assume a "rosette" habit under SD, e.g. henbane (*Hyoscyamus niger*). Runner formation in strawberry (*Fragaria*) plants occurs only under LD. The strawberry has a rosette habit which is not affected by daylength, but the axillaries have extended internodes (thus forming runners) under LD.

Tuber formation is also markedly affected by daylength, and is favoured by SD, as in the Jerusalem artichoke (*Helianthus tuberosus*) and in many wild species of potato (e.g. *Solanum andigena*). (The cultivated European varieties of potato are not very sensitive to daylength, and they can form tubers even under LD.) The onion (*Allium cepa*), on the other hand, requires LD for bulb-formation.

Daylength has a marked effect on the growth and leaf-fall of many wooded plants, as will be described below.

Sensitivity to Daylength Conditions

Many plants are extremely sensitive to changes in daylength as short as 15–20 minutes, so that the flowering time of some, such as certain varieties of rice (*Oryza sativa*), may be profoundly affected by even the relatively small seasonal changes in daylength found in the tropics. Similar sensitivity is shown by cocklebur (*Xanthium strumarium*) which only flowers under daylengths of $15\frac{3}{4}$ hours or less. Evidently the mechanism whereby such plants detect changes in daylengths is very sensitive.

Variation in sensitivity to daylength is shown in the number of photoperiodic cycles required to induce flowering. Thus, plants of *Xanthium* require exposure to only one SD cycle for flowering, and once induced to flower in this way they will continue to produce flowers for a further 12 months. The LD grass *Lolium italicum* will also flower in response to one LD cycle. However, these are extreme cases, and the majority of photoperiodic plants require more than one cycle. Moreover, even in *Xanthium*, the rate of floral development is more rapid after exposure to 2 or 3 SDs. In soybeans, the number of nodes at which flowers are found increases linearly with the number of SD cycles, up to at least 7. In many species, the daylength requirements vary considerably with the age of the plant, and frequently the minimum number of inductive cycles is found to decrease with age. Moreover, some SDP, such as soybeans, which will flower only under SDs when they are young, ultimately become capable of flowering even under LD.

LIGHT AND DARK PROCESSES IN SHORT-DAY PLANTS

A considerable amount of work has been carried out to learn more about the light and dark responses of SDP species, and our knowledge of this type of plant is considerably greater than for LDP.

It is important to ascertain, first, which part of the plant is concerned with the "detection" of daylength conditions. Clearly, in the transition from the vegetative to the flowering condition, the response is at the shoot apices, but it does not follow that "perception" of daylength conditions occurs in these parts of the plant. Indeed, it was shown by the Russian worker, Chailachjan, that the responses of SDP, such as *Chrysanthemum*, are determined by the daylength condition to which the *leaves* are exposed and that the shoot-apices appear to be relatively insensitive to daylength conditions. He was able to show this by exposing the leaves and shoot-apical regions independently to LD or SD conditions (Fig. 9.3); he found that it was only when the mature leaves were maintained under SD that flowering occurred. Thus, detection of the daylength conditions is effected by the leaves, although the response occurs at the shoot-apex. Chailachjan was quick to see that these observations imply that some "signal" must be transmitted from the leaves, which causes a response in the apices. We shall discuss the possible nature of this "signal" in a later section (p. 236).

In most plants the peak of photoperiodic sensitivity in the leaf appears to be reached when it has just attained its maximum size. At this stage quite small amounts of leaf tissue are sufficient to bring about flowering. Thus, in *Xanthium* 2 cm² are sufficient to bring about flowering under SD.

Although leaf tissue is the most sensitive to daylength conditions, the stem tissues of

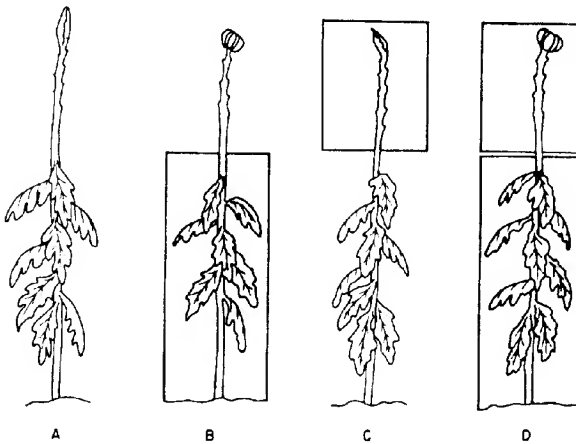


FIG. 9.3. Flowering occurs in *Chrysanthemum* when the *leaves* are exposed to short days (B and D), irrespective of whether the shoot apex is exposed to long days (B) or short days (D).

some species also show some sensitivity, a good example being seen in *Plumbago indica*, a SDP. Indeed, isolated internode sections of this species may be induced to flower under SD conditions in aseptic culture (Fig. 9.4), whereas under LD they remain vegetative.

The next problem which arises is whether the responses of SDP are determined by (1) the length of the daily light period, (2) the length of the dark period, or (3) the relative lengths of the light and dark periods. Now it can easily be shown that the responses of SDP are not controlled primarily by the total duration of light which they receive each day. This is shown by the fact that soybeans will flower if exposed to three cycles consisting of 12 hours light/12 hours dark, but will not do so if exposed to 36 hours of light, followed by 36 hours of dark, or if exposed to cycles consisting of 6 hours of light alternating with 6 hours of



FIG. 9.4. Induction of flowering *in vitro*. Flower of *Plumbago indica* L. "Angkor" produced on a segment of stem internode (7 mm in length) excised from a vegetative plant and planted aseptically on nutrient agar. Flower formation occurs only if the culture is placed under short days (10 hours of light). Under long days (16 hours of light), vegetative buds are produced. (Experiment by J. P. and C. Nitsch. Photo. Mlle B. Norreel, kindly supplied by Dr. J. P. Nitsch.)

lark (Fig. 9.10), although the total hours of light and dark are the same in all three régimes.

If we conduct experiments based upon the natural 24-hour cycles, it is impossible to vary the lengths of the light and dark periods independently; this can be done, however, if we use artificial sources of illumination, such as fluorescent lamps, since we can then choose any combination of light and dark periods we wish. K. C. Hamner was the first to take advantage of this fact in a series of experiments with *Xanthium* and soybean, which have now become "classical". Using soybean Hamner first investigated the effect of varying the length of the dark period, keeping a constant duration of light period of either (a) 4 hours, or (b) 16 hours. The results (Fig. 9.5) showed quite clearly that soybeans will not flower until the length of the daily dark periods exceeds about 10 hours, with either 4-hour or 16-hour photoperiods. Thus, the *critical dark period* for soybeans is about 10 hours, but the maximum flowering response is reached with dark periods of 16–20 hours. In similar experiments with *Xanthium*, Hamner showed that this species has a critical dark period of $\frac{1}{4}$ hours to flower.

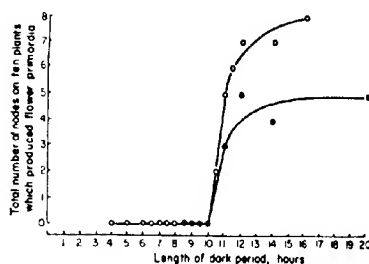


FIG. 9.5. Effect of various lengths of dark period, in association with constant 16-hour (—○—) or 4-hour (—●—) photoperiods, on flowering in soybean (*Glycine max*). (From K. C. Hamner, *Bot. Gaz.* 101, 658, 1940.)

Since SDP require a certain minimum period of darkness for flowering it may be asked whether they flower most readily in continuous darkness and whether they require any light. It is found, however, that although certain species, especially those with a storage organ such as a tuberous rootstock, will flower in continuous darkness, other species, such as soybeans, will not do so, but require a regular alternation of light and dark.

Having investigated the dark requirements of soybeans, Hamner then proceeded to study their light requirements. Using a constant dark period of 16 hours, he varied the length of the light period and found that the flowering response increased as the length of the daily light period was increased to about 12 hours (Fig. 9.6), but with longer light periods fewer flowers were formed and there was no flowering when the light period reached 20 hours, even though these light periods were associated with long (16-hour) dark periods. Thus, the conditions for flowering in soybean are (1) that the daily dark period must exceed 10 hours and (2) that the length of the light must not exceed a certain duration.

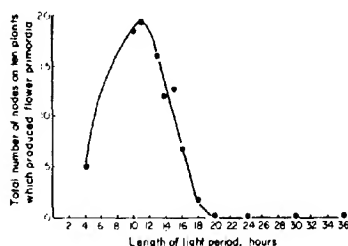


FIG. 9.6. Effect of varying length of photoperiod, with constant daily dark period of 16 hours, on flowering of soybeans. (From K. C. Hamner, *Bot. Gaz.* **101**, 658, 1940.)

Of course, under natural conditions, dark periods of 10 hours or more can only be accompanied by light periods of 14 hours or less, and in nature the flowering response is likely to be controlled by the length of the dark period, rather than the duration of the light period. For this reason, it would be more appropriate to refer to short-day plants as “long-night” plants. In some SDP, e.g. *Xanthium*, long photoperiods are not inhibitory to flowering, which is determined solely by the length of the dark period.

Having thus established that flowering in short-day plants is favoured by an alternation of light and dark, the question arises as to whether the light must precede the dark or vice versa. By means of ingenious experiments with *Xanthium*, which, as we have seen, requires only one SD for flowering, Hamner was able to show that a long dark period must be preceded by an adequate period of high-intensity light, and that a period of high-intensity illumination following the dark period also promotes flowering.

The Nature of the High-intensity Light Reaction

The light requirements of SDP during the photoperiod have been investigated quantitatively by Hamner and others. In general, it appears that the light requirements are relatively high. Thus the flowering response in soybeans increases as the period of exposure to daylight is increased from 1 to 8 hours. Moreover, with photoperiods of constant duration (5 to 10 hours), the number of flowers increases steadily with increasing light intensity.

These relatively high light requirements suggest that the process involved during the main photoperiod is photosynthesis. This conclusion is confirmed by the following facts: (1) carbon dioxide is necessary during the photoperiod, if flowering is to occur; (2) if sugar is sprayed on the leaves, then certain shortday plants can flower in complete darkness. Thus, there seems little doubt that the requirement for light in SDP is for adequate photosynthesis. This requirement might be expected, since an adequate supply of energy in the form of carbohydrates will clearly be necessary to permit the other metabolic processes, more specifically related to flowering, to occur in the leaves.

The "Dark" Reactions of Short-day plants

As we have seen, SDP require to be exposed to daily cycles which include a certain minimum period of darkness, the so-called "critical" dark period. This critical dark period appears to be relatively constant and independent of the length of the photoperiods over quite a wide range of the latter. At first sight it might be thought that the dark period plays a passive role, in the sense that the effects of darkness are simply due to the absence of light effects. Thus, it might be postulated that light has an inhibitory effect upon flowering, and that the flower-promoting effects of darkness are primarily due to the absence of such inhibitory effects. However, there are several pieces of evidence which suggest that certain positive flower-promoting processes occur during the dark period. Thus, it is known that the effectiveness of the dark period increases with temperature within certain limits, suggesting the occurrence of flower-promoting processes with positive temperature coefficients during the dark period.

A very remarkable feature of the dark period is that it must be *uninterrupted* if it is to be effective in promoting flowering—interruption by only a few minutes of light during the dark period may completely nullify its effect, so that flowering is inhibited. Thus, with *Xanthium*, 1 minute of light at an intensity of 150 foot-candles (fc) (≈ 1500 lux) during a 9-hour dark period suppresses flowering. This effect has been investigated in some detail in *Xanthium*, in which it was found that a "night-break" was most effective (maximum inhibition of flowering) when given 8 hours after the beginning of the dark period, regardless of the length of the latter, at least over the range 10–20 hours (Fig. 9.7). Thus we have the apparently paradoxical situation that light during the main photoperiod preceding a long dark period promotes flowering, whereas light given during the dark period inhibits

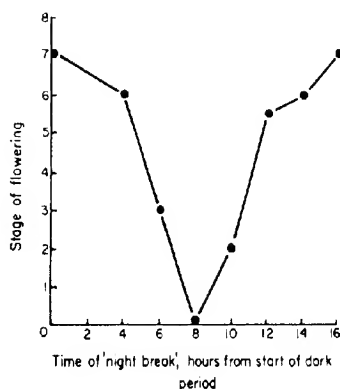


FIG. 9.7. Effect of night interruption, given at various times during dark period, on flowering in *Xanthium*. (From F. Salisbury and J. Bonner, *Plant Physiol.* 31, 141, 1956.)

flowering. We have seen that the requirement for light in the main photoperiod is probably for photosynthesis, to allow the formation of photosynthates necessary for metabolism of the leaf. However, it is now clear that the effects of a short interruption ("night-break") during a long dark period are not primarily due to photosynthesis but involve phytochrome. Detailed action spectra were determined for the night-break effect in soybeans and in *Xanthium* by placing seedlings with single leaves in different parts of the spectrum produced by a large spectrograph, and these action spectra proved to be almost identical to those which were later demonstrated for other phytochrome-controlled responses, such as seed germination and "de-etiolation" of shoots (p. 187). Moreover, R/FR reversibility was demonstrated for the night-break effect (Table 9.2), thus showing conclusively that this is a phytochrome-controlled response.

TABLE 9.2. Effect of daily interruptions of the dark period with several consecutive irradiations with red and far-red in sequence on flower initiation of *Xanthium* and Soybean

Treatment	Mean stage of floral development in <i>Xanthium</i> †	Mean no. of flowering nodes in <i>Biloxi</i> soybean
Dark control	6.0	4.0
R	0.0	0.0
R, FR	5.6	1.6
R, FR, R	0.0	0.0
R, FR, R, FR	4.2	1.0
R, FR, R, FR, R	0.0	—
R, FR, R, FR, R, FR	2.4	0.6

† See p. 200 for method of scoring.

Although flowering in the majority of species studied is inhibited by a relatively short night-break of a few minutes, other species require a longer night-break, e.g. the garden chrysanthemum requires a night-break of 4 hours.

The "escape-time" also varies considerably among different species; for example, in *Xanthium* the effect of R can be reversed if the application of FR is delayed for up to 40 minutes whereas in *Chenopodium album*, *Pharbitis nil* and *Kalanchoë blossfeldiana* FR must be given almost simultaneously after R to be effective.

The effects of R and FR given as night-breaks clearly indicate that the flower-promoting effects of a long inductive dark period are profoundly affected by the levels of P_{fr} in the leaf. At the end of the main photoperiod a high proportion of the phytochrome will be present in the P_{fr} form but, as we have seen, the levels of P_{fr} decline fairly rapidly during darkness, due to degradation or reversion. However, if a short exposure to R light is given after several hours of darkness high levels of P_{fr} will be restored, with inhibitory effects on the flower-promoting processes (Fig. 9.8), which apparently require low P_{fr} levels.

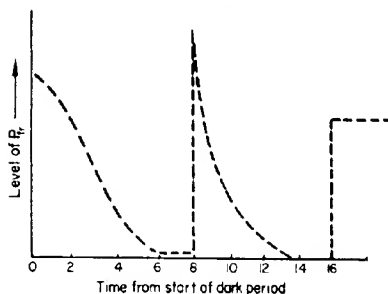


FIG. 9.8. Hypothetical changes in the P_{fr} form of phytochrome in leaves during a dark period which has been interrupted by a short exposure ("night-break") to red light.

Although, with short night-breaks FR reverses the inhibitory effect of R, with longer periods (e.g. 60 minutes) FR acts in the same manner as R, and is inhibitory to flowering, especially after short photoperiods. This effect seems to indicate that although the low values of P_{fr} caused by FR are not effective over short periods, they can be effective if maintained over longer periods.

Flower Promoting Effects of P_{fr} in Short-day Plants

There is evidence that although high levels of P_{fr} are inhibitory to flowering when they occur during a long inductive dark period, as a result of a night-break, at other times of the daily cycle flowering may be promoted by high P_{fr} levels. Thus, flowering in seedlings of *Pharbitis nil* is reduced by FR (which will reduce the level of P_{fr}) given at the end of the main photoperiod. It would appear, therefore, that at that stage in the daily cycle flowering is favoured by high P_{fr} levels, whereas after several hours of darkness it is inhibited, as indicated by the effects of a R night-break. Thus, flowering in SDP is apparently promoted by high P_{fr} at the end of the main photoperiod, and inhibited by high P_{fr} later during the dark period, indicating that phytochrome has a dual action in the photoperiodic control of flowering.

RESPONSES OF LDP

As in SDP, the effect of a long dark period on LDP is nullified by a short night-break, and hence *promotes* flowering in plants of this type.

Studies on the action spectrum for the night-break effect in barley (*Hordeum*) and *Hyoscyamus* indicated, once again that it is the red region (660 nm) which is effective and that the effects of R are reversed by FR in some, but not all, LD species. However, LDP are

less sensitive to night-breaks than SDP, and although a night-break of 1–2 hours is effective in some LDP, the effect is quantitative and the maximum effect may require still longer periods.

When supplementary illumination is used to extend a short photoperiod given as daylight, it is found that a mixture of R and FR is more effective in promoting flowering in LDP than R alone (Fig. 9.9). The R and FR need not be given simultaneously. Thus, in *Lolium temulentum*, when R alone was used to extend 8 hours of daylight it was ineffective in inducing flowering, but if this R was interrupted by a few hours of FR, flowering was induced. It was found that there was an optimum time for exposure to FR, and it appears that following the end of an 8-hour photoperiod of sunlight, high P_{fr} levels (caused by R)

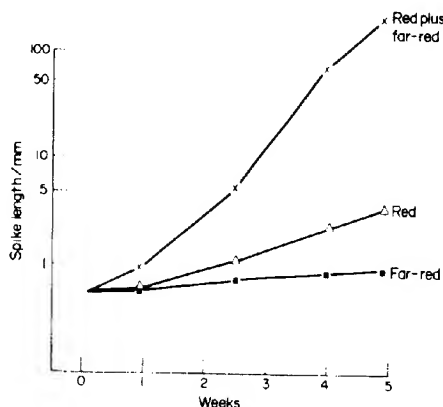


FIG. 9.9. Effect of red, far-red and mixed red plus far-red light on flowering in *Lolium temulentum*. Plants received 8 hours of daylight plus a further 8 hours of supplementary light of restricted wavelengths. (From D. Vince-Prue, *Photoperiodism in Plants*, McGraw-Hill, London, 1975.)

reduce the flowering response, whereas later in the dark period high P_{fr} tends to favour flowering and at this time R promotes. FR appears to promote flowering from about the sixth hour from the beginning of the daily photoperiod, but after about the eighteenth hour FR has little effect and flowering is now promoted by R. However, a diurnal change in P_{fr} level is not essential for LDP as it is in SDP. Thus, it appears that, as in SDP, high P_{fr} promotes flowering at certain times and inhibits at others, but the sequence of promotion and inhibition is opposite in LDP and SDP. Thus, we see that there are close analogies between the behaviour of SDP and LDP, but in certain respects their phytochrome-controlled responses are opposite, so that long dark periods promote flowering in SDP but inhibit it in LDP (Fig. 9.10). In both types of plant, moreover, the effect of the dark period is nullified by a night-break which results in high P_{fr} levels. During the earlier part of the

daily cycle, flowering in SDP is promoted by high P_{fr} levels, so that they require a sequence of high P_{fr} during the main photoperiod and low P_{fr} during an ensuing long dark period (Fig. 9.11). By contrast, LDP do not require such a diurnal fluctuation in P_{fr} , since they flower even in continuous light (under incandescent lamps which produce low P_{fr} values), but under suitable experimental conditions a diurnal fluctuation in sensitivity to P_{fr} levels can be demonstrated.

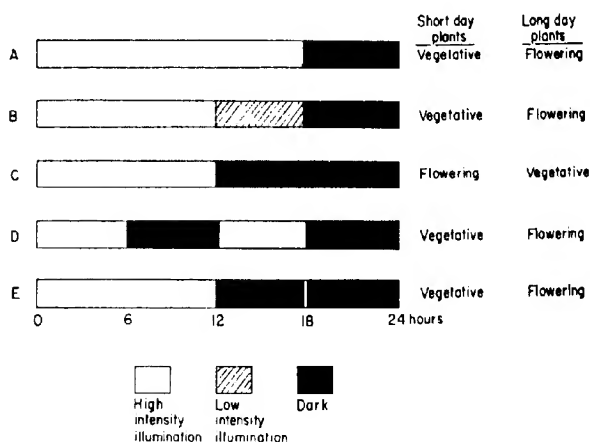


FIG. 9.10. Summary of responses of short-day plants and long-day plants to various photoperiodic régimes.

THE FLOWERING STIMULUS

We have seen that the response of the plant to daylength depends upon the conditions to which the leaves are exposed, although the response occurs at the shoot meristem. This observation immediately suggests that under favourable daylength conditions some flower-promoting "stimulus" is formed in the leaves and conveyed from there to the meristems. This hypothesis was put forward by Chailachjan very shortly after his discovery that the perceptive organs in photoperiodism are the leaves, and he postulated that a flower hormone (which he called "florigen") is synthesized in the leaves under favourable daylength conditions and transmitted to the growing points. As we shall now see, there is a great deal of evidence to support this hypothesis, but some 40 years after Chailachjan first postulated the existence of "florigen", it still remains to be isolated and characterized chemically. A considerable number of highly interesting experiments on the transmission of the flower hormone have been carried out and these will now be described.

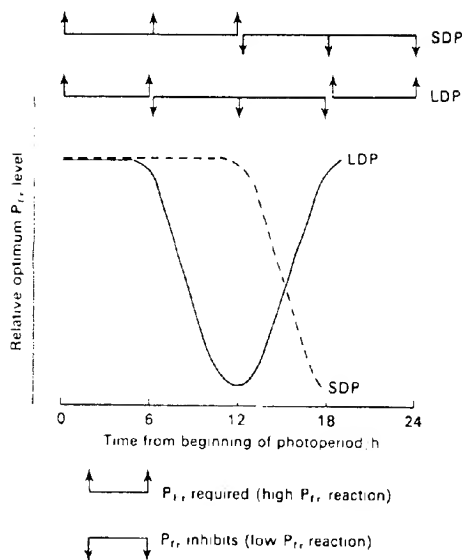


FIG. 9.11. Schematic representation of the "high P_{fr} " and "low P_{fr} " reactions in the flowering of long-day plants and short-day plants. In the SDP the P_{fr} -requiring process occurs early in the photoperiod and is followed by a period when P_{fr} inhibits flowering. In LDP similar reactions are seen, but their timing is markedly different. (From D. Vince-Prue, *Photoperiodism in Plants*, McGraw Hill, London, 1975.)

Hamner confirmed Chailachjan's experiments, using *Xanthium*. He showed that flowering of *Xanthium* plants under SD does not occur if all the leaves are removed, but only one-eighth of one leaf is sufficient to result in flowering (Fig. 9.12). In experiments with "two-shoot" plants of *Xanthium*, it was found that the stimulus arising from a single leaf is sufficient to cause flowering not only in the shoot on which it is borne, but also on the second shoot from which all leaves have been removed.

A large number of experiments have shown that the flowering-stimulus can be transmitted across a graft union. Thus, Hamner grafted together the stems of pair of plants of *Xanthium*: when one of the plants of a pair was exposed to SD, not only did this plant flower, but also its grafted partner which had not been exposed to SD (Fig. 9.12). In further experiments plants of *Xanthium* were exposed to SD, their tops were then decapitated and scions from other *Xanthium* plants which had been maintained under LD were grafted on to them. In due course, it was found that the "LD" scions flowered, although they had not themselves ever been exposed to SD.

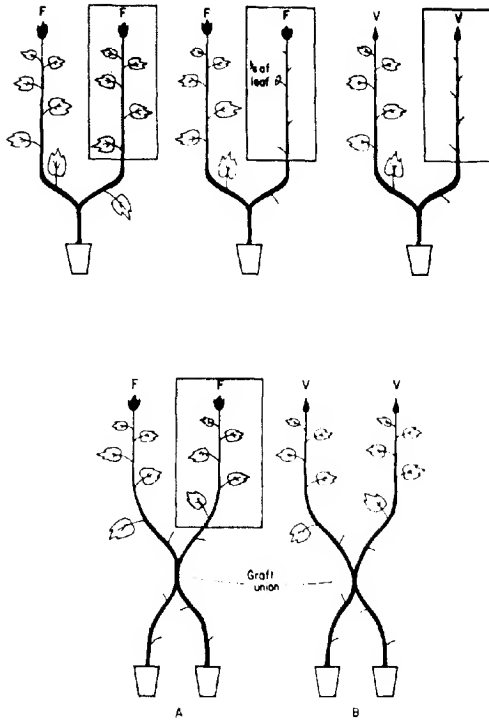


FIG. 9.12. Above: Two-branched plants of *Xanthium*, one branch of which was exposed to short days, the rest of the plant being exposed to long days. Both branches have flowered, provided that the "short-day" shoot has at least one-eighth of a leaf, but not if it is completely defoliated (right).

Below: A. Two *Xanthium* plants were approach grafted, and the top of one plant was exposed to short days while the other plant was maintained under long days. Both plants have flowered.

B. Both plants exposed to long days; neither have flowered. (From K. C. Hammer, *Cold Spring Harbor Symp.* 10, 49, 1942.)

In still more striking experiments carried out with various species, including *Xanthium* and *Perilla*, single leaves were removed from "donor" plants which had been exposed to SD and grafted on to plants maintained under LD. In due course the LD "receptor" plants flowered, showing that the stimulus may be transmitted from a single leaf.

Grafting experiments with LDP have been very much fewer than with SDP, but the results have proved essentially the same.

Experiments of the type just described leave little doubt that some "stimulus", presumably a substance of a hormonal nature, is formed in the leaves under favourable daylength conditions and is transmitted through the stem to the apical meristems.

Although the flower hormone has not yet been isolated from SDP, we know quite a lot about the time required for its synthesis in the leaf and about its transport in the plant. If *Xanthium* plants, each with a single leaf, are exposed to one SD cycle, and the leaves are removed at various times after the end of the long dark period, it is found that flowering does not occur unless they are allowed to remain on the plant for at least 2–4 hours following the end of the dark period, but greater flowering responses are obtained if the leaves are left for 1–2 days before removal, indicating that movement out of the leaf is a slow process. There seems no doubt that transport of the hormone occurs through living tissues, presumably by the phloem, since transport is stopped by steaming a zone of stem between the leaf and the apical region, but it may be transported through other living tissues.

Attempts have been made to estimate the rate of transport of the hormone. One method was to use two-shoot plants of *Pharbitis* in which a single donor leaf on one branch was exposed to SD, and the differences in time of flower initiation on the second branch (kept under LD) were determined for varying distances between the donor leaf and the receptor bud. Methods of this type, which admittedly are very indirect, suggest that the rate of movement of the flower hormone is much slower (2–4 mm/hour) than the normal rate of translocation of sugars in the phloem. An experiment with *Pharbitis* has given evidence of much more rapid transport, however (Fig. 9.13). Evidence was obtained that the flowering stimulus had moved over a distance of 102 cm in 2 hours, i.e. a rate of 51 cm/hour, which is commensurate with the rate of movement of photosynthates in the phloem.

PHOTOPERIODIC "AFTER-EFFECT" AND THE INDUCTION OF LEAVES

As we have seen, a short-day plant does not have to be maintained continuously in short days in order to flower. After a certain number of favourable photoperiodic cycles a SD plant will flower even though it may subsequently be transferred to LD. Thus, as we have seen, *Xanthium* will ultimately flower in response to a single SD cycle, although a considerable number of LD cycles may intervene between the SD and the ultimate appearance of flowers. Thus, certain species showed a marked photoperiodic "after-effect", and a plant is said to become "induced" after exposure to SD.

This after-effect of favourable photoperiodic cycles is evidently a property of the leaves, which apparently continue to produce "flower-hormone" even after they have been transferred from favourable to unfavourable daylength conditions. Thus, the inductive response of the whole plant reflects the inductive changes occurring in the leaves. This conclusion is supported by many experiments. For example, Lona subjected a single leaf of a *Perilla* plant to SD, the other leaves remaining under LD conditions. After this period of SD

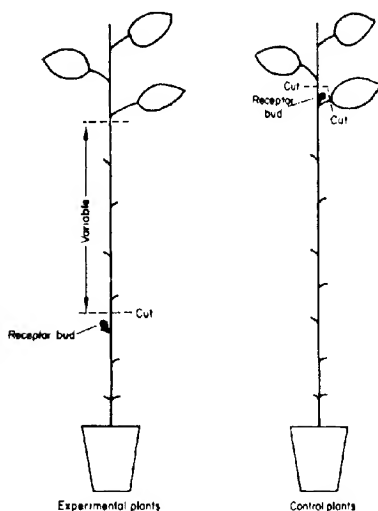


FIG. 9.13. *Pharbitis* plants of various heights were decapitated above the uppermost fully expanded leaf and all leaves except for the three upper ones were removed. The plants were divided into two groups. In the experimental group all axillary buds were removed except for one receptor bud at the base of the stem, and in the control group all buds were removed except for the one in the axil of the lowest donor leaf. The plants were given a single dark period which ranged in duration from 14 to 16 hours in different batches of plants, and at the end of the dark period all leaves and stems were cut off just above the receptor buds. No flower buds were formed in either group when the leaves were removed 14 hours after the start of the dark period. Flower buds were, however, initiated in many plants whose donor leaves were removed 16 hours after the start of the dark period, irrespective of the length of stem between the donor leaf and the receptor bud. The tallest plant had a stem 102 cm long between the lowest donor leaf and the receptor bud at the base, and initiated two flowers. As there was no flowering in the lowest donor leaf in control plants given a 14-hour dark period, the flowering stimulus must have moved through 102 cm of stem within the last 2 hours of the 16-hour dark period, yielding a rate of movement of more than 51 cm/hour. (From G. Takeba and A. Takimoto, *Bot. Mag., Tokyo*, 79, 811–14, 1966.)

treatment, the leaf was allowed to remain under LD for 4 weeks, when it was removed and grafted on to a vegetative *Perilla* plant, which in due course flowered. Thus, the leaf “remembered” the previous SD treatment although 4 weeks of LD treatment intervened between the end of the last SD cycle and the grafting on to the receptor plant. What is the basis of this after-effect shown by the leaves?

Two theories have been put forward to account for the after-effect. Chailachjan postulated that a store of flower hormone is accumulated in the leaf under favourable conditions and is gradually exported from it for a long period even under favourable conditions. On

the other hand, another Russian worker, Moshkov, postulated that the metabolism of the leaf somehow becomes *permanently* changed in response to favourable daylength conditions, so that it continues actively to produce flower hormone even if it is subsequently transferred to unfavourable daylengths. The available evidence seems strongly to support Moshkov's theory. Thus, it would seem unlikely that in the experiment of Lona described above, sufficient flower hormone would still remain in the SD-treated leaf after 4 weeks of LD treatment to induce flowering when it was grafted on to a vegetative plant.

Even stronger evidence that leaves may become permanently changed under favourable daylength conditions is provided by certain experiments of Zeevaart. In one experiment, *Perilla* plants were first exposed to forty SD cycles and the leaves were then grafted on to vegetative plants growing under LD. Every 14 days, ten of these leaves were removed and regrafted on to a new group of vegetative plants. It was found that the leaves continued to induce flowering in each new group of vegetative stocks on to which they were grafted (Fig. 9.4). There was no diminution in the flowering response even with the fifth group of leaves, which had been maintained under LDs for 10 weeks. In a second experiment there was no diminution in the flower-inducing effect after seven such graftings, although more than 3 months had elapsed since the leaves had received the SD treatment. Thus, in *Perilla*, the leaves appear to become permanently induced.

It is found that the state of induction is strictly localized within the *Perilla* plant and even within the individual leaves. Thus, if single pairs of leaves of *Perilla* are exposed to a series of SD cycles and the rest of the plant is kept under LD, it is found that only those leaves which directly received SDs are capable of inducing the plants to flower when grafted on to them (Fig. 9.14). In other experiments by Lona, only one half of single leaves were exposed to SD, and the other half of each leaf to LD. The leaves were then divided longitudinally and grafted separately on to vegetative receptor plants. Only those half leaves which had been directly exposed to SD were capable of inducing flowering.

The situation is different in *Xanthium*, however. If a single leaf from a flowering *Xanthium* plant (A) is grafted on to a vegetative plant (B) growing under LD, this will flower, as has already been described. If other young leaves, which have never themselves been exposed to SD, are taken from plant B and grafted on to further vegetative plants (C) these will also flower. Thus, the leaves of plant B have themselves become induced by the grafting of an induced leaf from plant A, although these leaves of B have never directly been exposed to SD. This effect is described as *secondary induction*, and it probably explains why *Xanthium* does not revert to the vegetative condition when it is transferred from SD to LD conditions, since all the new leaves formed will become secondarily induced. On the other hand, the new leaves formed by a *Perilla* plant which is transferred from SD to LD will not be induced and they appear to inhibit the flower-promoting effects of the older leaves which became induced by the previous SD treatment (see below), and hence the plant reverts to the vegetative condition.

Although *Xanthium* and *Perilla* show marked persistence of photoperiodic induction of leaves, it is not known how general are these effects. Reversion of whole plants to the vegetative condition on exposing to a minimal number of SDs and then returning them to LD

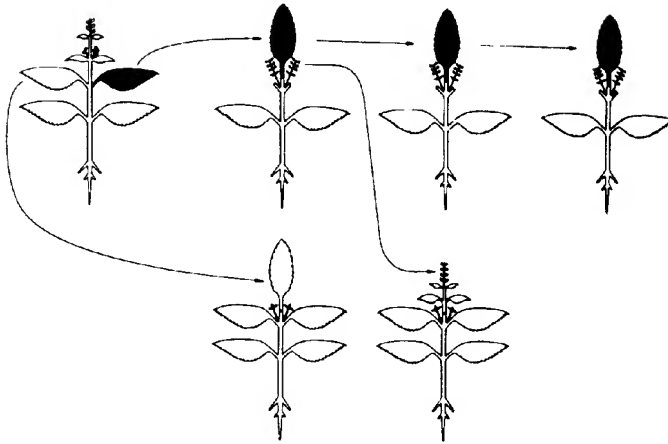


FIG. 9.14. Permanency of the photo-induced state in *Perilla* (see text). Black: leaves exposed to short days; in outline: plants on long days. Leaves which have directly received SD treatment will induce flowering when grafted successively to negative receptor plants, whereas non-induced leaves from the original or from the receptor plants, and flowering shoots from the latter, will not do so. (From A. Lang, *Encycl. Plant Physiol.* 13(1), 1416, 1965, after data from Zeevaart.)

certainly occurs in soybean and a number of other species, but whether this is due to "de-induction" of the leaves exposed to SD, or to the production of new, non-induced leaves, as in *Perilla*, is not known.

As Zeevaart has pointed out, the permanent induction of leaves makes it necessary to distinguish between two distinct phenomena, viz.

- (1) the *induced state* (i.e. the ability to produce the floral stimulus), gradually built up under the influence of favourable daylengths, which is irreversible and strictly localized in some plants;
- (2) the *floral stimulus*, which is transmissible from induced leaves to the growing points where it exerts its morphogenetic effect.

NATURAL FLOWER-INHIBITING EFFECTS

Several lines of evidence suggest that not only flower-promoting processes, but also flower-inhibiting effects occur in photoperiodic plants. Thus, if one branch of a two-shoot plant of soybean or *Perilla* is exposed to SD and the other to LD, the latter will not flower

unless its own leaves are removed. That is to say, the flowering stimulus from the "donor" branch does not produce any effect in the "receptor" branch in the presence of LD leaves. Similarly, if a scion from a vegetative plant of *Perilla* is grafted on to a stock which has previously been maintained under SD, the scion will only flower if its leaves are removed. Thus, in some SD species, LD leaves exert an inhibitory effect on flowering.

This inhibitory effect of LD leaves is only manifested if they occur between the shoot apex and the source of the flowering stimulus. Thus, if a single leaf of *Kalanchoë* is exposed to SD and the remainder of the plant is maintained under LD, the presence of LD leaves between the SD leaf and the shoot apex prevents flowering, but if all leaves are removed above the SD leaf and only LD leaves below are allowed to remain the plant will flower. A LD leaf is particularly inhibitory if it is immediately above the SD leaf (i.e. on the same orthostichy). LD leaves on the opposite side of the stem from the SD leaf have less inhibitory effect.

It has been suggested that these inhibitory effects of non-induced leaves can be interpreted in terms of interference with the translocation of the flower hormone, which, it is assumed, is carried with the main stream of photosynthates. The leaves which supply the greater part of the photosynthates to the shoot apical region are the uppermost mature leaves. If the latter are SD leaves, then they will supply flower hormone, as well as photosynthates, to the shoot apex, but if LD leaves are interposed between the apex and the SD leaves, then the supply of both photosynthates and flower hormone from the SD leaves to the apex will be reduced.

Other authors have argued that these inhibitory effects of leaves can only be interpreted in terms of specific flower-inhibitory substances; that is to say, they postulate that there are flower-inhibitory hormones as well as flower-promoting ones, but this conclusion is based upon indirect evidence.

Other flower-inhibiting effects are seen in experiments in which SDP such as *Kalanchoë* are exposed to SD cycles between which are intercalated one or more LD cycles. It can be shown that if the plants are exposed alternately to two SD cycles and one LD cycle, flowering is completely inhibited, and a careful analysis of the data indicates that the effect of the intercalated LD cycle is not merely "neutral" but positively inhibitory to flowering. In this type of experiment it would seem that we are dealing with inhibitory effects of LDs within the leaf itself, and not with the export of a flower-inhibitor to the shoot apex.

Flower-inhibiting effects also occur in LDP, since, as we have seen, in these plants long dark periods appear to have an inhibitory effect. It seems likely, however, that the mechanism of inhibition in this case is different from that occurring in the leaves of SDP when they are kept under LD.

It might be asked whether, if there are flower-inhibiting processes in both SDP and LDP, it is necessary also to postulate active flower-promoting processes? Might not the flowering of SDP when transferred to SD conditions be due primarily to the removal of the flower-inhibitory processes occurring under LD? If so, we may not need to postulate the existence of the elusive "flower hormone". However, most workers in this field consider that other experimental evidence points strongly to the existence of both flower-promoting and

flower-inhibiting processes. Thus, the observation that a single leaf taken from a flowering plant of *Xanthium* will induce flowering in a vegetative plant maintained under LD is difficult to explain in terms simply of the removal of flower-inhibition in the receptor plant. It would seem, therefore, that the regulation of flowering in both SDP and LDP may involve the interplay of both flower-promoting and flower-inhibiting processes.

TIME MEASUREMENT IN PHOTOPERIODISM

We have seen that flowering in SDP occurs when they are kept under daylength conditions in which the length of the night exceeds the critical dark period and that some plants can detect differences in the length of the dark period of as little as 15 minutes. Thus, *Xanthium* has a critical dark period of $8\frac{1}{2}$ hours at 25°C . A difference of only 15 minutes in the length of the dark period can determine whether or not sugar cane will flower. It is clear, therefore, that these species have rather accurate time-measuring mechanisms. Several suggestions have been made as to the nature of this mechanism.

Thus, the "clock" might be of the "hour-glass" type, in which the time taken for a particular substance to accumulate or be depleted to a certain threshold value may be the time-measuring process. Since flowering in SDP requires that the phytochrome in the leaves shall be in the P_r form, and that there is a gradual conversion of P_{tr} to P_r during the first hours of darkness (p. 210), it has been suggested that the critical dark period may represent the time taken for P_{tr} to decline to a certain level. However, it is found that the reaction $P_{tr} \rightarrow P_r$ is effectively complete after 2–3 hours of darkness, whereas the critical dark period of *Xanthium* is $8\frac{1}{2}$ hours.

Again, if the length of the critical dark period is determined by the time taken for the conversion of P_{tr} to P_r , then irradiation with far-red light at the beginning of the dark period should greatly reduce the length of the critical dark period for flowering in SDP, by hastening the conversion of P_{tr} to P_r , but this is found not to be the case for all but one of the species so tested. On these and other grounds it seems unlikely that the rate of conversion of P_{tr} to P_r is the factor determining the length of the critical dark period of SDP. We are therefore forced to consider other hypotheses regarding the time-measuring mechanism, and among these is the "Endogenous Rhythm" hypothesis of Bünning.

ENDOGENOUS RHYTHMS IN PHOTOPERIODISM

It is many years since Bünning first drew attention to the existence of persistent rhythms in plants. He investigated the diurnal movement of leaves in the runner bean (*Phaseolus multiflorus*) in which the primary leaves rise during the early part of the day and later fall towards evening and reach a minimum position during the night; they then start rising again towards the morning. These movements are regulated by turgor changes in the "pulvini" of the leaves.

Now, if a bean seedling is exposed to a period of daylight and then kept in continuous darkness for several days, it will continue to show the typical diurnal movements, rising and falling on a 24-hour cycle even though it is itself not being exposed to the natural alternation of light and dark. That is to say, the bean plant shows a persistent endogenous rhythm in its leaf movements. If the position of the leaf is plotted against time then we obtain a sinusoidal curve. As the period of darkness is extended, the amplitude of the movements gradually declines to zero. The plant then needs a further exposure to light to set the rhythm in train again.

Since these early observations on leaf movements, it has been shown that many other processes in plants show a regular diurnal rhythmicity, including the following:

- (1) Opening and closing of flowers.
- (2) Root pressure.
- (3) Growth rate.
- (4) Respiration and other metabolic processes.
- (5) Activity of certain enzymes.
- (6) Mitosis and size of nucleus.
- (7) Discharge of fungal spores, e.g., *Pilobolus*.

The length of a single cycle of an endogenous rhythm when "free-running" in darkness is approximately, but not exactly, 24 hours. It is important that under natural conditions the endogenous rhythm of the plant should be synchronized with the diurnal cycle of day and night, and this is achieved by the *entrainment* of the endogenous rhythm by the natural day and night. Although the rhythms are endogenous in the sense that, once started, they can be maintained for several days even in continuous darkness, nevertheless, many such rhythms require the stimulus of the change from dark to light or vice versa, to start them off. Thus, synchronization between endogenous rhythm and natural day and night is achieved by daily setting of the rhythm by a "light-on" signal at dawn, or a "light-off" signal at dusk.

Thus, there can be no doubt as to the existence of endogenous rhythms in plants. Now Bünning postulated that there is also an endogenous rhythm in photoperiodic sensitivity, and he put forward a theory of photoperiodism based upon this endogenous rhythm. He suggested that during the day SDP are in a "photophile" phase, when light is favourable for flowering, and that during the night they are in a "skotophile" phase, when darkness is favourable and light is inhibitory to flowering. Thus we have to envisage a regular rhythm in photoperiodic sensitivity, as the plants enter first the photophile and then the skotophile phase. It was postulated that SDP will flower when the daily light and dark periods correspond with the photophile and skotophile phases of the plants, i.e. when they are under SD. Under LD light will extend into the skotophile phase and will inhibit flowering. The responses of LDP have proved more difficult to account for on this theory, and these responses may be the reverse of those of SDP.

Bünning's theory has been subjected to certain criticisms and put to various experimental tests, some of which have not given the results predicted by the theory. Nevertheless, there

now seems no doubt that there is an endogenous rhythm in photoperiodic sensitivity in at least some species. This rhythm has been demonstrated in various ways.

Thus, Hamner grew soybean plants on a wide range of different cycle lengths; each plant received 8 hours of light followed by a dark period the length of which varied from 8 to 62 hours. It was found that the plants flowered maximally when the total cycle length was 24 hours or a whole multiple thereof, i.e. 48 hours, 72 hours, but remained vegetative when the cycle length amounted to about 36 or 60 hours (Fig. 9.15). These results strongly

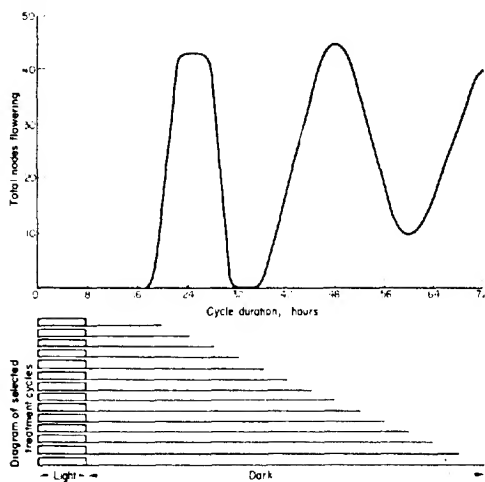


FIG. 9.15. Flowering responses of Biloxi soybeans to cycles by various lengths. Plants were exposed to seven cycles, each cycle consisting of 8 hours of high-intensity light (1000-1500 fc) and associated dark periods of various lengths. Total nodes flowering per 10 plants is plotted against cycle length. (After K. C. Hamner, Chapter 13 in *Environmental Control of Plant Growth*, Academic Press, New York, 1963.)

suggest that the light and dark processes in soybean are geared to a 24-hour cycle, so that when the cycle length is 24, 48 or 72 hours in length, the endogenous "photophile" and "skotophile" phases will correspond with the environmental light and dark periods, but that when the latter are on 36- or 60-hour cycles they will be out of phase with the endogenous rhythm of the plants, which will therefore not flower. It is also found that the vegetative growth of tomato plants is best when the cycle length is 24 hours or a whole multiple thereof, but is poor on cycle lengths of 36 or 60 hours. However, other species, including *Kalanchoë* and *Xanthium*, show no marked responses to variations in cycle length.

A second type of approach to the problem of endogenous rhythms in photoperiodism has involved the introduction of short light interruptions ("night-breaks") at different

stages of long dark periods, to determine whether there is any evidence of rhythmic variations in photoperiodic sensitivity, as judged by the flowering response. Experiments have been carried out with plants maintained on 48-hour or 72-hour cycles in which, following a short photoperiod (of say 10 hours), night-breaks are given at various stages during the following 38- or 62-hour dark periods. In soybean and in *Chenopodium rubrum*, such night-breaks are found to be inhibitory to flowering during the early and late part of a 38-hour dark period, corresponding to the skotophile phases postulated by Bünning. With 62-hour dark periods three inhibitory phases are found, corresponding again to the predicted three skotophile phases in a 72-hour cycle (Fig. 9.16).

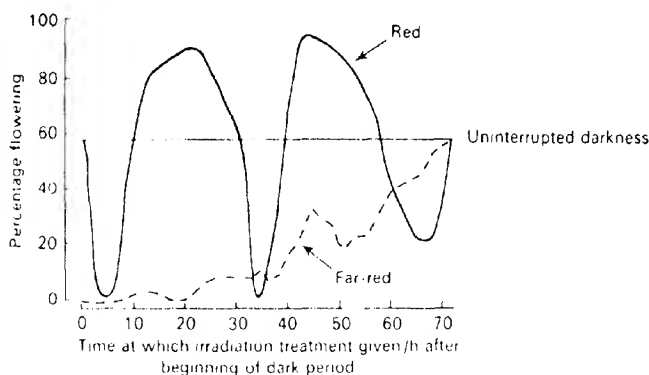


FIG. 9.16. Flowering of *Chenopodium rubrum* with a single 72-hour dark period interrupted at different times by 4 minutes of red light or by 10 seconds of intense far-red light. (From D. Vince-Prue (1975) adapted from B. G. Cumming, S. B. Hendricks and H. A. Borthwick, *Can. J. Bot.* **43**, 825, 1965.)

Thus, two different types of approach have given results, with certain species, which appear to indicate that there is, indeed, an endogenous rhythm in photoperiodic sensitivity, similar to that postulated originally by Bünning. However, other species have given different results. Thus, although "night-breaks" given during long dark periods show rhythmicity in *Kalanchoë*, flowering in this species is little affected by the cycle length. Again, night-break experiments with *Pharbitis nil* show very little evidence of rhythmicity, and the experimental data can be interpreted in terms of an "hour-glass" model.

Bünning originally postulated that the photoperiodic cycle is set off by the onset of dawn, which starts the oscillation, and that the skotophile phase follows on 12 hours after this "light-on" signal. However, later experimental data seem to indicate that in some species the oscillation is established by the transfer from light to dark, i.e. by a "light-off" signal. There is some evidence that phytochrome may be involved in the "light-off" response. For example, in *Lemna perpusilla*, an endogenous rhythm in carbon dioxide

output is established by a light-off signal at the end of a period of red light, and R/FR reversibility has been demonstrated for the light phase.

From the evidence presented there seems little doubt that some plants do show an endogenous periodicity in photoperiodic sensitivity. It still remains an open question, however, whether such rhythms occur in all species showing photoperiodic responses. Thus, endogenous rhythms may modify the photoperiodic responses in some species but may not be a universal factor in the photoperiodism of all plant species. Moreover, the demonstration of rhythmicity by the type of experiment described above does not imply the correctness of Bünning's hypothesis regarding photophile and skotophile phases.

It is clear that the occurrence of endogenous rhythms implies the existence of some sort of "oscillator" mechanism in the plant, but the nature of this oscillator is still completely unknown. However, such an oscillator could serve as a time-measuring mechanism or physiological clock, but whether this is the actual time-measuring mechanism involved in photoperiodism must remain an open question for the present.

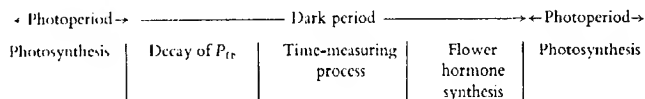
THE SEQUENCE OF PROCESSES LEADING TO HORMONE SYNTHESIS

We have seen that it has been possible to recognize a number of partial processes leading to the production of the flowering stimulus in SDP, and we are now in a position to summarize briefly the present state of knowledge regarding the sequence and interrelations of these processes.

Firstly, there is a requirement for a high-intensity light reaction (photosynthesis) which is met during the photoperiod, and which evidently provides the energy and substrates necessary for the processes occurring during the ensuing dark period. However, there is reason to believe that phytochrome effects are also involved during the photoperiod, since the flowering of SDP, such as *Kalanchoë blossfeldiana*, when maintained in continuous darkness for long periods, is promoted by a short daily period of red light, suggesting that a minimal level of P_{fr} during the photoperiod is required for flowering. Normally, at the end of the photoperiod, the phytochrome will be present in approximately equal concentrations of P_r and P_{fr} . During the first few hours of the dark period, P_{fr} decays, so that P_r now predominates. Flower hormone synthesis does not commence until a certain minimum period of darkness (critical dark period) has elapsed and it then proceeds rather rapidly in the next few hours. It is apparently necessary for phytochrome to be present in the P_r form in order for flower hormone synthesis to proceed, since we know that a short interruption with red light (which converts P_r to P_{fr}) inhibits flowering. However, the commencement of hormone synthesis is apparently not directly controlled by the decay of P_{fr} , since the duration of the critical dark period is apparently considerably longer than the time required for P_{fr} decay. Hence it has been suggested that there must be some "time-measuring" mechanism, which more directly controls the commencement of hormone synthesis. We have seen that it is possible that the time-measuring mechanism involves an endogenous

rhythm in certain unknown processes, and that the role of phytochrome in flowering may lie in its effects on the time-measuring processes, rather than directly on the process of flower hormone synthesis.

These ideas are summarized as follows:



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The Physiology of Flowering— II. Temperature and Other Factors

VERNALIZATION

As we have seen, photoperiodic responses to seasonal variation in daylength conditions will account for the periodicity in flowering behaviour of many plant species, both temperate and tropical. It will be noticed, however, that among the examples of temperate species which show photoperiodic responses there were relatively few spring-flowering species, although it is a matter of everyday observation that there is a considerable number of "flowers that bloom in the spring", and many of these spring-flowering forms, such as celandine (*Ficaria verna*), primroses (*Primula vulgaris*), violets (*Viola* spp.), etc., show marked seasonal behaviour, and remain vegetative for the remainder of the year after the spring flush of flowers. It might have been expected that spring-flowering is a response to the short days of winter, but this does not appear to be the case in many species.

Daylength is not, of course, the only environmental factor showing an annual variation, and clearly temperature also shows well-marked seasonal changes, especially in temperate regions, although there is considerable variations from day to day and from year to year. We now know that seasonal variations in temperature, as well as in daylength, have a profound effect upon the flowering behaviour of many species of plant.

The clue to the importance of temperature as a regulator of flowering came first from the studies of Gassner in 1918 on the flowering of cultivated varieties of cereals. Cereals such as wheat (*Triticum*) and rye (*Secale*) can be grouped into two classes depending upon whether they are to be sown in the autumn ("winter" varieties) or in the spring ("spring" varieties). Winter wheat sown in the autumn, or spring wheat sown in the spring, both flower and mature in the following summer. If the sowing of the winter wheat is delayed until the following spring, however, it fails to ear and remains vegetative throughout the growing season. It seems unlikely that the necessity for sowing winter wheat in the autumn is simply to secure a longer growing season as such, since autumn-sown plants make relatively little growth during the winter, and winter wheat plants from spring sowings certainly seem to make adequate leaf development, yet they will not flower.

Gassner therefore investigated the effects of different temperature régimes during the germination and early growth of winter and spring rye. He sowed winter and spring rye in sand at different dates between 10th January and 3rd July and kept them at the following temperatures during germination: 1–2°C, 5–6°C, 12°C and 24°C. They were later planted out-of-doors. He found that the temperature during germination had no influence on the subsequent flowering of spring rye, and all seedlings planted out on the same date flowered at approximately the same time, irrespective of the temperature treatment during germination. In winter rye, however, only the plants which had been germinated at 1–2°C flowered regardless of the planting date out-of-doors. Seedlings that were germinated at temperatures above 1–2°C were found to flower only if they were planted not later than March or early April, so that they would have been exposed to some actual chilling out-of-doors (in Central European climatic conditions). Gassner concluded that whereas the temperature conditions during early growth do not affect the flowering of spring rye, the flowering of winter rye depends on its passing through a cold period, either during germination or later (Fig. 10.1).

This work of Gassner was later followed up in the U.S.S.R. and a great deal of work was carried out there, particularly in relation to possible economic applications. The severity of the Russian winter in many regions does not permit autumn sowing of winter wheats, which, however, usually have a higher yield than spring varieties. A technique was devised by Lysenko, whereby the cold treatment required by winter wheat was applied to



FIG. 10.1. Effect of vernalization of flowering in "Perkus" winter rye (*Secale cereale*). Left: maintained for several weeks at 1°C after germination; right: seed unvernallized. (From O. N. Purvis, *Ann. Bot.* 48, 919, 1934.)

the seed before sowing in the spring. The method used was to allow partial soaking of the seed, so that there was sufficient imbibition of water to allow slight germination and growth of the embryo, but not sufficient for complete germination. Seed of winter wheat in this condition, which was exposed to cold treatment by burying in the snow, attained flowering and maturity in the same season if sown in the spring. The technique came to be known as *vernalization*, and this term has subsequently been extended to other treatments involving exposure to winter chillings, not only at the seed stage, but also at later stages of development of the plant.

Types of Plant showing Chilling Requirements for Flowering

We have seen that chilling of winter wheat is effective in stimulating subsequent flowering whether given during the early stages of germination or later, when considerable development of leaves has occurred. Later work has shown that many species have a chilling requirement for flowering, including winter annuals, biennials and perennial herbaceous plants. Winter annuals are species which normally germinate in the fall and flower in the early spring. They include such species as *Aira praecox*, *Erophila verna*, *Myosotis discolor* and *Veronica agrestis*.

It now appears that winter annuals and biennials are effectively monocarpic† plants which have a vernalization requirement—they remain vegetative during the first season of growth and flower the following spring or early summer, in response to a period of chilling received during the winter. The need for biennials to receive a period of chilling before flowering can occur has been demonstrated experimentally for a number of species, including beet (*Beta vulgaris*), celery (*Apium graveolens*), cabbages (and other cultivated forms of *Brassica*), Canterbury bells (*Campanula medium*), honesty (*Lunaria biennis*), foxglove (*Digitalis purpurea*) and others. If plants of foxglove which normally behave as biennials, flowering in the second year after germination, are maintained in a warm greenhouse they may remain vegetative for several years. In regions with a mild winter climate cabbages may grow for several years in the open without “bolting” (i.e. flowering) in the spring, as they do in areas with cold winters. Such species have an obligate requirement for vernalization, but there a number of other species in which flowering is hastened by chilling but will occur even in unvernallized plants; such species showing a facultative cold requirement include lettuce (*Lactuca sativa*), spinach (*Spinacia oleracea*) and late-flowering varieties of pea (*Pisum sativum*).

As well as biennial species, many perennial species show a chilling requirement and will not flower unless they are exposed each winter to cold conditions. Among common perennial plants which are known to have such a chilling requirement are primrose (*Primula vulgaris*), violets (*Viola* spp.), wallflowers (*Cheiranthus cheirii* and *C. allionii*), Brompton stocks (*Mathiola incana*), certain varieties of garden chrysanthemum (*Chrysanthemum*

† See p. 281 for an explanation of this term.

morifolium), Michaelmas daisies (*Aster* spp.), Sweet William (*Dianthus*), rye-grass (*Lolium perenne*). Perennial species require revernalizing every winter.

It seems very probable that many other spring-flowering perennials will prove to have a chilling requirement when they are investigated. Bulbous spring-flowering plants such as daffodil (*Narcissus*), hyacinth, bluebell (*Endymion nonscriptus*), crocus, etc., do not have chilling requirements for flower initiation, the flower primordia being laid down within the bulb during the previous summer, but their growth is markedly affected by temperature conditions. For example, in tulip flower initiation is favoured by relatively high temperatures (20 °C), but the optimum temperature for stem elongation and leaf growth is initially 8–9 °C, rising to 13 °C, 17 °C and 23 °C at successively later stages. Similar temperature responses are shown by hyacinth and daffodil (*Narcissus*).

In many species flower initiation does not occur during the chilling period, but only after the plants are exposed to higher temperatures following chilling. However, some plants, such as Brussels sprouts, have to remain at low temperatures until flower primordia have actually been formed. It appears that all the species with a chilling requirement for flowering can be vernalized at the "plant" stage, i.e. as leafy plants, but not all species can be vernalized at the "seed" stage, as can winter cereals. Among the other species which can be vernalized at the seed stage are mustard (*Sinapis alba*) and beet (*Beta*). On the other hand, cultivated varieties of *Brassica* (cabbages, Brussels sprouts) and celery cannot be vernalized in the seed stage but the seedlings must attain a certain minimum size before they become sensitive to chilling, and thus show a "juvenile" phase. Generally, those species which can be vernalized at the seed stage are facultative cold-requiring plants, whereas those which can only be vernalized at the plant stage show an obligate chilling requirement.

Many plants with a chilling requirement resemble long-day plants, in that they have a rosette habit in the vegetative phase, and show marked internode elongation associated with flowering.

The requirement for chilling for flower initiation must not be confused with a chilling requirement for the removal of bud dormancy (see p. 259). Thus, many woody plants flower in the spring, but the flower initials are laid down within the bud during the preceding summer and they have a chilling requirement, not for flower initiation, but for the removal of bud dormancy.

Species showing both Chilling and Photoperiodic Responses

Interactions between vernalization and photoperiodic responses have been studied in a number of species. Henbane (*Hyoscyamus niger*) exists in annual and biennial forms, corresponding to the spring and winter forms of rye. The annual form does not require vernalization, but is a long-day plant, which flowers in the summer. The biennial form requires vernalization, followed by long days, for flowering to take place.

As an example of a perennial plant which shows both vernalization and photoperiodic responses, we may mention perennial rye-grass (*Lolium perenne*). In this species, flowers are

initiated in response to winter chilling, but long days are required for emergence of the inflorescence, so that elongation of the flowering stem does not commence until the day-length exceeds 12 hours in March. The new tillers (lateral shoots) which emerge during the spring and summer are unvernized and remain vegetative throughout the growing season, until the following winter. Consequently, flowering of perennial rye-grass is seasonal and restricted to the spring and early summer.

Vernalization requirements are less common among short-day plants, but the garden chrysanthemum constitutes an example which has been studied intensively. Certain varieties of chrysanthemum have a vernalization requirement which must be met before they will respond to short days. After the parent plant has flowered in the autumn a number of horizontally growing rhizomes emerge from the base of the plant and grow just beneath the surface of the soil. Under outdoor conditions these new shoots will become vernalized by natural winter chilling, and in the spring they grow into normal, upright leafy shoots, which grow vegetatively under the long days of summer and initiate flowers in response to short days in the autumn. If, however, the plants are not exposed to chilling during the winter, but are grown under warm conditions in a greenhouse, the new shoots do not become vernalized, and although they grow actively throughout the summer, they are incapable of forming flowers in the short days of autumn. Thus, the chrysanthemum provides another example in which the new shoots arising each year need to be vernalized, since the vernalized condition is not transmitted from the parent shoots, as it is in rye (p. 232).

Genetical aspects of vernalization responses have been studied in several species. In rye, the difference between spring and winter forms is found to be controlled by a single major gene, the requirement for vernalization of the winter forms being recessive to the non-requirement of spring rye. The opposite situation exists in henbane, where the biennial habit (vernalization required) is dominant to the annual habit (no chilling requirement). In other species, however, the inheritance of chilling responses is more complex and in rye-grass (*Lolium perenne*) several genes appear to be involved.

Physiological Aspects of Vernalization

Intensive studies on the physiological changes underlying vernalization have been carried out on relatively few species and our knowledge of the subject is based largely on the work of Gregory and Purvis on winter rye, of Melchers and Lang on henbane (*Hyoscyamus niger*) and of Wellensick on several other species.

The work of Gregory and Purvis has established a number of important characteristics of the processes occurring during vernalization of rye. Firstly, they showed that the changes occur in the embryo itself and not in the endosperm, as had been suggested. This was done by removing the embryos from the grain and cultivating them on a sterile medium containing sugar; such embryos were given a period of cold treatment and showed the typical hastened flowering responses given by vernalized grain, when planted. It was even possible

to vernalize isolated shoot apices, which had been removed from the embryos and cultivated under sterile conditions. Such apices developed roots and regenerated into seedlings which ultimately flowered in response to the earlier chilling treatment. Moreover, it was shown that vernalization may even be effected whilst the young embryos are still developing in the ear of the mother plant. This was done by enclosing the developing ears in vacuum flasks packed with ice, or by removing developing ears and keeping them in a refrigerator until mature. In this way it was shown that vernalization is effective in embryos, even if commenced only 5 days after fertilization.

It has been shown that in older plants it is the shoot apical region which must be chilled. This was shown for celery, beet and chrysanthemum by placing a cooling coil around the shoot apical region. Thus, whereas in photoperiodism it is the *leaves* which are the sensitive organs to daylength, in vernalization the shoot apex itself is sensitive to the chilling temperatures. Wellensiek has shown that in *Lunaria* young leaves are also capable of being vernalized, but older leaves, which have ceased growth, do not respond, unless they show some cell division at the base of the petiole. Wellensiek maintains that only tissues which have dividing cells are capable of being vernalized.

For the majority of species, the most effective temperatures are just above freezing, viz. $1-2^{\circ}\text{C}$, but temperatures ranging from -1°C to 9°C are almost equally effective. Hence freezing of the cells is not necessary to bring about the changes occurring during vernalization, suggesting that physiological, rather than purely physical processes are involved. This conclusion is confirmed by the observation that cold treatment of rye grain is ineffective under anaerobic conditions, indicating that probably aerobic respiration is essential. By cultivating excised embryos on media with and without sugar it was shown that a supply of carbohydrate is necessary during cold treatment. Thus, although the metabolism of most plants is considerably retarded at cool temperatures, there seems little doubt that vernalization involves active physiological processes, but the nature of these processes is still quite unknown.

It is found that the degree of hastening of flowering in rye varies with the duration of exposure to cold, the longer the cold treatment the shorter the period from sowing to flowering, up to a certain limit, beyond which further cold treatment has no further hastening effect (Fig. 10.2A). Quite short exposures to cold, such as 7–11 days, have a noticeable vernalizing effect, and this effect increases progressively with the duration of treatment.

The vernalization process can be reversed by exposing the grains to relatively high temperatures ($25-40^{\circ}\text{C}$) for periods of up to 4 days. Seeds treated in this way show a reduced flowering response, and are said to be "devernalized". As the period of chilling increases, it becomes increasingly difficult to reverse the effect, and when vernalization is complete, high temperature is ineffective. Devernalized grains can again be vernalized by a further period of chilling.

Once the rye plant has been vernalized it appears that the condition is transmitted to all new tissues formed subsequently, so that all new laterals are also vernalized. Indeed, if the main shoot apex is removed, so that the laterals are stimulated and then these are decapitated to stimulate secondary laterals and so on, it is found that even fourth-order laterals

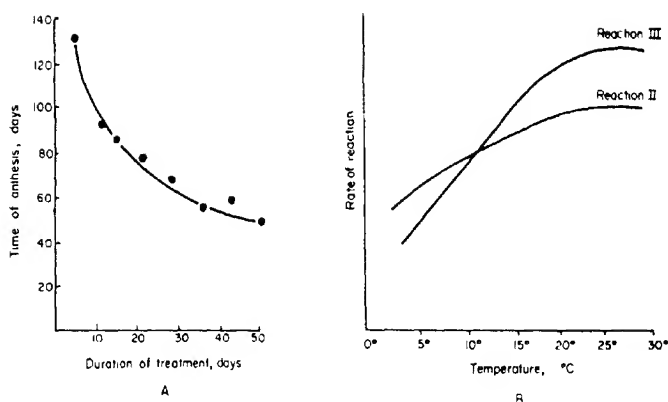


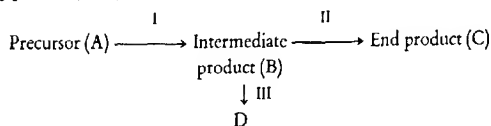
FIG. 10.2. A. Effect of the duration of chilling of seed, on subsequent flowering behaviour of Petkus winter rye. Curve indicates time to anthesis from planting, for various periods of vernalization. (From O. N. Purvis and F. G. Gregory, *Ann. Bot.* 1, n.s. 569, 1937.)

B. Hypothetical scheme, illustrating two reactions with different temperature coefficients (see text).

are still fully vernalized although the apices of these laterals were not present at the time of the chilling treatment. Thus, the vernalized condition is transmitted from a parent cell to its daughter cells in cell division and it does not appear to be "diluted" in the process.

The Nature of the Changes occurring during Vernalization

One of the striking features of vernalization is that, at first sight, it appears to involve processes which go on more rapidly at lower than at higher temperatures. This effect is most unusual for chemical processes and yet we must assume that the changes occurring during vernalization are essentially enzyme-controlled reactions showing the usual characteristics of such reactions. How then are we to explain the apparent "negative temperature coefficient" of vernalization? One very simple hypothesis which has been put forward postulates the occurrence of two separate processes competing for a common substrate, each having positive (though different) temperature coefficients:



In this scheme reaction II and III compete for the common "Intermediate product" B. Suppose reaction III has a higher temperature coefficient than reactions I and II (Fig. 10.2B).

This means that high temperatures will favour reaction III and more of B will be diverted into this reaction, so that little C will be formed. When, however, the temperature is markedly reduced, this will reduce the rate of reaction III more than that of II (since by definition reaction III shows greater response to changes in temperature). Consequently, reaction II will be favoured at the reduced temperature and C will accumulate. C will, therefore, be formed at lower temperatures but not at higher temperatures. Thus, the overall production of C will appear to have a "negative temperature coefficient", although each of the three involved reactions has a positive temperature coefficient. There is no direct evidence to support this hypothesis, but it is of value as indicating how an overall process may proceed more rapidly at lower temperatures, without contravening the normal laws of chemical reactions.

We have already seen that vernalization appears to involve relatively stable changes, so that once the fully vernalized state has been attained by meristematic tissue, it appears to be transmitted by cell-lineage without "dilution". This conclusion in turn may imply that the vernalized state is transmitted through some self-replicating cytoplasmic organelle, but it is equally possible that certain genes become activated during vernalization and that once this has occurred, this change is transmitted to the daughter nuclei during division.

The Flowering Stimulus in Vernalization

Although so much study has been devoted to vernalization, our understanding of the nature of the physiological and biochemical processes involved is still very fragmentary. It is not yet clear whether a specific transmissible flower hormone is formed as a result of vernalization, although there is evidence that this may be the case, at least in some species. We have seen that in photoperiodism some of the strongest evidence for the existence of a flower hormone is provided by grafting experiments, and somewhat similar results have been obtained with certain species showing a vernalization requirement.

A notable example of flower induction by grafting in a vernalized plant is provided by henbane. If a leaf from a vernalized plant of the biennial variety of henbane is grafted on to an unvernallized stock of the same variety, the latter is induced to flower without chilling. A similar response can be obtained by grafting on to the unvernallized biennial variety a leaf from any of the following: (1) the annual variety of henbane (a LDP, with no vernalization requirement); (2) *Petunia hybrida*, an annual LDP; (3) tobacco, day-neutral variety (4) tobacco, variety "Maryland Mammoth" (SDP), under either SD or LD. Thus, transmission of a flowering stimulus can take place between cold-requiring and non-cold-requiring plants, even of different genera. Similar results have been obtained in the biennial species beet (*Beta vulgaris*), cabbage (*Brassica oleracea*), carrot (*Daucus carota*) and *Lunaria biennis*. On the basis of these results, Melchers and Lang suggested that a transmissible flowering stimulus, which they called *vernalin*, is formed as a result of chilling in biennial plants.

On the other hand, in some species, including *Chrysanthemum*, it has not proved possible

to obtain transmission of a flowering stimulus from vernalized to non-vernalized plants by grafting. Moreover, if the tip region of the plant was given localized cold-treatment and hence flowered, the other buds not directly chilled remained vegetative. Similarly, when the tip of an unvernallized radish plant was grafted on to a vernalized one, flowering did not occur. These latter results are consistent with the conclusion that the *vernalized condition* (as opposed to a flowering stimulus) is only transmitted through cell division, i.e. by "cell lineage". It would seem, therefore, that we must make a distinction between the vernalized ("thermo-induced") state and the formation of a flowering hormone, just as we saw that in photoperiodism we have to distinguish between the induced state of a leaf, and the transmissible stimulus formed in it.

The question then remains as to whether there is a specific flowering stimulus, vernalin, formed in plants with a chilling requirement. In most of the successful grafting experiments, referred to above, the donors were from species which have a requirement for both chilling and LDs or for LDs alone, and the experiments were usually carried out under LD conditions. Moreover, it is generally found necessary that the donor shoots should have leaves. Thus, it seems possible that "vernalin" is identical with the flower hormone produced in the leaves of LDP. Melchers and Lang have argued against this last conclusion, on the ground that leaves of Maryland Mammoth tobacco will induce flowering in stocks of biennial henbane, both under SD and under LD, although "Maryland Mammoth" itself will not flower under the latter conditions (Fig. 10.3). On the other hand, biennial *Hyoscyamus* causes flower formation in Maryland Mammoth tobacco only if the donor has been vernalized. These results suggest that there is a distinct transmissible agent which in cold-requiring plants arises only after chilling, whereas in non-cold-requiring ones its formation is not dependent on photoperiodic induction, and hence it cannot be identical

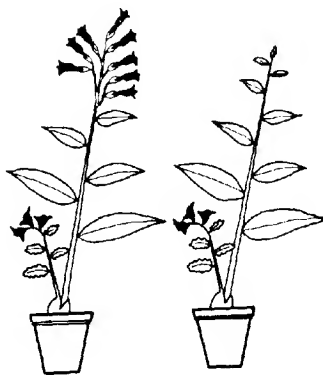
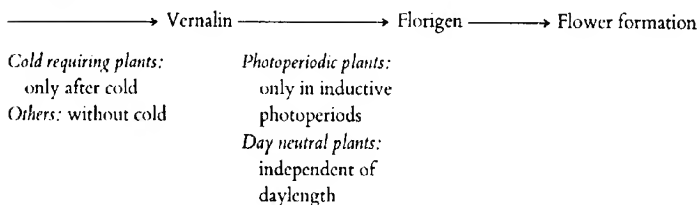


FIG. 10.3. Flower induction in non-vernalized biennial *Hyoscyamus niger* by Maryland Mammoth tobacco donors in short days (left) and long days (right). Note the flowering response caused by the non-photoinduced donor. (From Lang, 1965—see Further Reading.)

with florigen. It has therefore been suggested that vernalin and florigen are interdependent, and that vernalin must be present if florigen is to be formed. Thus, the sequence† of events for cold-requiring plants may be summarized as follows:

Low temperature → vernalized state → vernalin → florigen.

More generally, the sequence† can be written as follows:



THE NATURE OF THE FLOWERING STIMULUS

Attempts to Isolate the Flowering Stimulus

As we have seen, the evidence from the various types of grafting experiment described in Chapter 9 strongly suggests that a flowering stimulus arises in the leaves of both SDP and LDP under favourable daylength conditions and is transported to the meristems where it causes a vegetative apex to change to the flowering condition. Moreover, it would seem that the same type of flowering stimulus is produced in both SDP and LDP, as the following experiment suggests. *Kalanchoë blossfeldiana* is a SDP, whereas the species of *Sedum* *ellacombianum* and *S. spectabile* of the same family (Crassulaceae) are LDP. If vegetative shoots of *Sedum* are taken from plants growing under SD and grafted on to *Kalanchoë* under SD, not only do the stocks of the latter flower, but also the *Sedum*, which will not itself flower under SD (Fig. 10.4). Thus, a stimulus is produced in the SDP *Kalanchoë* which will cause flowering in the LDP *Sedum*. Conversely, if vegetative shoots of *Kalanchoë* are taken from plants growing under LD and grafted on to *Sedum* plants under these conditions, again both stock and scion will flower. Thus, the LDP *Sedum* produces a flowering stimulus under LD which is capable of causing the SDP *Kalanchoë* to flower. These experiments suggest that the flowering stimulus is identical in both SDP and LDP.

A range of experimental results of the type described are consistent with the hypothesis that a specific flower hormone arises in the leaves under inductive daylength conditions, and is transmissible by grafting. As we have seen, grafting experiments with species having a chilling requirement have also given evidence of a transmissible flowering stimulus.

Although the circumstantial evidence for the existence of flower hormones is very

† From Lang, 1965 (see Further Reading).

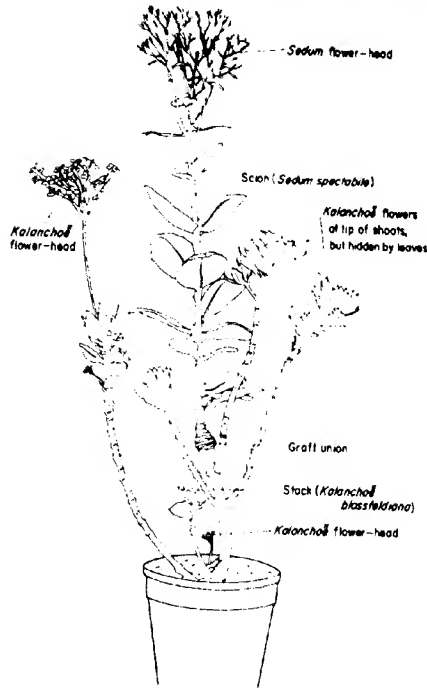


FIG. 10.4. Transmission of flowering stimulus from short-day plant, *Kalanchoe blossfeldiana*, to long-day plant, *Sedum spectabile*, by grafting. A scion from a vegetative plant of *Sedum* growing under short days was grafted on to a flowering plant of *Kalanchoe*, also under short days. The scion of *Sedum*, as well as the *Kalanchoe* stock, has flowered under these conditions. (Original by I. D. J. Phillips from plant supplied by J. Hillman.)

strong, repeated attempts to extract a specific flower hormone from SDP over the past 30 years have nearly always given negative results. There have been certain claims that extracts of *Xanthium* plants can promote flowering, but this claim has not always been repeatable by other workers. On the other hand, flower-promoting, gibberellin-like substances have been extracted from LDP (p. 239).

This failure to isolate the flower hormone from SDP is baffling, but there are several possible explanations, the three most likely ones being as follows:

- (1) The hypothetical hormone may be present in extremely low concentrations, or it may be very unstable, so that existing methods of extraction and detection are not sufficiently sensitive.

- (2) Since the naturally occurring growth hormones are relatively simple compounds, it has generally been assumed that the hypothetical flower hormone will also prove to be a fairly simple substance. There is no reason, however, to exclude the possibility that the flower hormone has a complex, unstable molecule, such as protein or nucleic acid. The fact that the stimulus can be transmitted by grafting, but cannot be extracted, is consistent with this hypothesis, since special techniques would be required to isolate it. The same difficulties of extraction are encountered with certain plant viruses. Attempts to detect differences between the proteins of flowering and non-flowering plants of *Xanthium* or other species have given no clearcut results. Studies are now being carried out to determine whether there are qualitative differences in specific RNAs between flowering and non-flowering plants.
- (3) There may be no specific flower hormone—the induction of flowering may depend rather upon *quantitative* differences in the level of certain well-known substances occurring in plant tissues.

We have already seen that dramatic morphogenetic changes can be induced in tissue cultures by varying the relative levels of substances such as auxins and kinins in the culture medium (p. 148). Thus, it is argued, the transition from the vegetative to the flowering condition may be controlled by changes in the levels of, for example, certain growth hormones such as auxins, gibberellins and kinins, or by the balance between these hormones. We shall now consider briefly this possibility.

Effects of Gibberellins and Other Growth Hormones on Flowering

Although attempts to regulate flowering through application of auxins have almost all been unsuccessful (apart from the exceptional case of the pineapple, p. 135) it has been found that a number of LDP can be induced to flower under SD by application of GA_3 (Table 10.1). LDP which respond to GA_3 are typically species which form a pronounced rosette under SD and which show marked internode elongation ("bolting") under LD. When GA_3 is applied to such species growing under SD there is a very marked stimulation of internode elongation and this process is accompanied by flower initiation.

Gibberellin is also effective in stimulating flowering the "long-short-day" plants *Bryophyllum crenatum* and *B. daigremontianum*, which normally require to be exposed first to LD and then to SD for flower initiation but which will flower under continuous SD if treated with GA_3 . Thus, GA_3 substitutes for the LD requirement in these species.

A number of species which normally require vernalizing before flower-initiation will occur can also be induced to flower by external application of GA_3 (Table 10.1, Fig. 10.5). Thus, in these species GA_3 apparently replaces the chilling requirement. However, GA_3 will apparently not stimulate flowering of unvernallized rye and certain other species, although it will stimulate stem elongation in these species. In general, treatment of seeds

TABLE 10.1. Species showing flowering under non-inductive conditions in response to applied gibberellic acid (GA_3)

A. Long-day Plants	
<i>Anagallis arvensis</i> (Pimpernel)	<i>Petunia hybrida</i> (Petunia)
<i>Cichorium endivia</i> (Chicory)	<i>Raphanus sativus</i> (Radish)
<i>Hyoscyamus niger</i> (Henbane, annual)	<i>Rudbeckia bicolor</i> (Coneflower)
<i>Lactuca sativa</i> (Lettuce)	<i>Silene armeria</i>
<i>Papaver somniferum</i> (Poppy)	<i>Spinacia oleracea</i> (Spinach)
B. Plants with Chilling Requirement	
<i>Apium graveolens</i> (Celery)	<i>Digitalis purpurea</i> (Foxglove)
<i>Avena sativa</i> (Oat)	<i>Hyoscyamus niger</i> (Henbane, biennial)
<i>Beta vulgaris</i> (Sugar-beet)	<i>Matthiola incana</i> (Stock)
<i>Bellis perennis</i> (Daisy)	<i>Myosotis alpestris</i> (Forget-me-not)
<i>Brassica oleracea</i> (Cabbage)	<i>Solidago virgaurea</i> (Golden rod)
<i>Daucus carota</i> (Carrot)	

with GA_3 is not effective in stimulating flowering, even in species which respond to seed vernalization.

These observations raise the question of whether the endogenous gibberellins are not the "flower hormone" in LDP and species showing vernalization responses. Thus, it might be postulated that in LDP growing under SD the level of endogenous gibberellins is too low for flowering, and that the effect of LD is to raise the level of endogenous gibberellins to the threshold necessary for flowering. Indeed it has been shown for certain species, including spinach and henbane, that there is a marked rise in the levels of endogenous gibberellin when the plants are transferred from SD to LD conditions. Moreover, extracts of gibberellins of the LDP *Rudbeckia*, growing under LD conditions, will induce flowering in plants of the same species growing under SD. Similarly, during vernalization of the biennial species, hollyhock (*Althaea rosea*), there is an increase in the levels of a gibberellin-like substance which will stimulate flowering in *Rudbeckia*, although it will not stimulate flowering in unvernallized hollyhock.

Although these latter observations suggest that changes in endogenous gibberellins may be important in flowering, there is other evidence against the hypothesis that flowering in LDP and plants which require vernalization is regulated primarily by gibberellins, including the following:

- (1) As we have seen, there is evidence that the flowering stimulus is identical in both LDP and SDP, and yet gibberellins are quite ineffective in inducing flowering in most SDP.
- (2) Not all LDP and species with a chilling requirement can be induced to flower in response to applied GA_3 . (However, certain of these latter species can be induced to flower when other types of gibberellin are applied. For example, GA_4 and GA_7 are effective in promoting flowering in *Myosotis alpestris*, which does not respond to



FIG. 10.5. Effect of vernalization and gibberellic acid on flowering of carrot. *Left*: untreated control plant; *right*: plant chilled for 8 weeks; *centre*: plant unchilled but treated with $10\ \mu\text{g}$ GA_3 per day. (From A. Lang, *Proc. Nat. Acad. Sci., U.S.A.* **43**, 709, 1957.)

GA_3 . Thus, in some species there appear to be rather precise requirements with respect to the nature of the gibberellin which will stimulate flowering, and these requirements differ between one species and another.)

- (3) Nearly all rosette plants respond to GA_3 by elongation of the internodes, including those species which do not flower in response to this treatment. Moreover, in henbane, when flowering is induced by LD treatment, the formation of flower primordia *precedes* the elongation of the internodes, whereas in response to GA_3 treatment plants of this species kept under SD begin to show internode elongation before the flower primordia appear. Again, stem growth in *Silene armeria* can be completely suppressed with AMO-1618 while flower formation proceeds normally, thus showing that

stem growth and flower formation are independent processes controlled by different hormones. Furthermore, the genetic analysis of *Silene* clearly shows that there are separate genes for stem growth and for flower formation. These facts suggest that internode elongation and flower initiation are distinct processes and that the primary effect of GA_3 is on internode elongation, with flower-initiation occurring as a secondary effect of stem elongation in certain species.

- (4) Exogenous GA_3 will not stimulate flowering of "normal" genotypes of red clover (*Trifolium pratense*), but in certain non-flowering genotypes of this species both LD and GA_3 are necessary for flowering. Thus, in these non-flowering genotypes, GA_3 does not substitute for LD, suggesting that some other flower-promoting factor is also normally involved in this species.

Thus, although there is good evidence that changes in the levels of endogenous gibberellins may play an important role in the flowering responses of LDP, it would seem that this is not "the whole story" in LDP and certainly the responses of SDP cannot be accounted for solely in these terms.

In addition to the stimulation of flowering by gibberellins, a number of other growth-regulating substances, both natural and unnatural, have been found to promote flowering in some species under certain conditions. Thus, flowering can be induced in pineapples not only by synthetic auxins (p. 135) but also by ethylene. Under certain conditions kinetin and adenine will promote flowering in *Perilla* and zeatin does so in the aquatic plant *Wolffia microscopica*. Similarly, the naturally occurring growth inhibitor, abscisic acid (p. 67) promotes flowering in *Pharbitis*, *Fragaria* and *Ribes*, while the synthetic growth retardants CCC and B.9 promote flowering in a number of species, including apple and pear trees. A number of other substances, including tri-iodobenzoic acid, maleic hydrazide, vitamin E and even sugars, have been reported to promote flowering in a few species.

Nevertheless, in the majority of SDP and in some LDP, flowering cannot be induced by any combination of the known naturally occurring hormones. Hence we cannot at present account for the flowering behaviour of the majority of species in terms of interaction between known growth hormones.

The conclusion that flowering behaviour cannot be accounted for in terms of the known growth hormones, and yet all attempts to extract a specific flower hormone have been unsuccessful, may indicate that we have adopted an oversimplified approach to the problem, in assuming that flowering is controlled by a single, specific hormone. The isolation and identification of the flowering stimulus remains one of the most challenging problems in developmental plant physiology.

SEX EXPRESSION AND GROWTH HORMONES

There is some evidence that growth hormones may be involved in the determination of sex in some plants. This has come from studies of dioecious species (male and female flowers

on separate plants) such as hemp (*Cannabis sativa*), and of those monoecious species in which the male and female organs, stamens and ovaries are borne in separate flowers on the same plant (e.g., some varieties of cucumber (*Cucumis sativus*)). Treatment of genetically male plants of hemp with an auxin spray causes female flowers to be produced. In monoecious cucumber varieties, it is normally the case that male flowers develop during the earlier stages of growth and that female flowers form only later on. However, application of an auxin to the leaves of young cucumber plants results in an acceleration of the transition from production of male flowers to production of female flowers. It has been suggested, therefore, that female flowers or female parts of flowers tend to differentiate under conditions of higher auxin concentration than do male flowers or parts. This conclusion is supported by the finding that genetically determined forms of cucumber which bear only male flowers contain lower levels of endogenous auxin than the normal hermaphrodite forms. Femaleness in cucumber is also enhanced by treatment with ethylene, or etrel (a commercial preparation of 2-chloroethane-phosphonic acid, which is converted to ethylene in plant tissues, p. 138).

Gibberellin treatment of monoecious cucumber plants, in contrast to auxin treatment, increases the number of male flowers formed. Treatment of gynoeceious cucumber (i.e. a dioecious variety which normally produces only female flowers) with gibberellin results in the formation of male as well as female flowers. Moreover, endogenous gibberellin levels are lower in the gynoeceious types than in the normal hermaphrodite forms. It is possible, therefore, that sex expression in plants is effected by a balance between endogenous auxins and gibberellins. The effect of auxin on flower sexuality may involve the participation of ethylene (see p. 65).

CHANGES OCCURRING IN THE SHOOT APEX DURING FLOWER INITIATION

We have seen that the transition from the vegetative to the reproductive condition involves drastic changes in the structure of the shoot apical meristems (p. 44). The earliest steps in this transition have been studied in a number of SDP and LDP. Indeed, SDPs such as *Xanthium* or *Chenopodium*, and LDPs such as *Lolium temulentum*, in which flowering may be induced by exposure to a single inductive cycle, provide very favourable material for studying the transition, since the timing of the latter can be rigorously controlled to within a few hours. Studies on *Xanthium* have revealed that the earliest changes leading to flower initiation can be first detected about 4 days after exposure to a single SD cycle, but when the plants are exposed to two SD cycles changes can already be detected at the end of this treatment. As we have seen (p. 44), the first changes occur in the region between the central mother cells and the rib meristem, and involve cell division in this region. A stage is soon reached in which a "mantle" of small, densely staining and actively dividing cells overlies a central core of more vacuolated cells (Fig. 2.18).

It is clear that when a plant changes from the vegetative to the flowering phase many genes must be brought into action, including those which control flower and fruit development. Thus, we must postulate some mechanism which regulates the "switching on" of the flowering genes. We know very little concerning the nature of such gene-switching mechanisms in higher plants, but this subject is discussed further in Chapter 13. Since genes, i.e. DNA, control development by the synthesis of specific enzymes, through a process involving various types of RNA, it is evident that nucleic acid metabolism is likely to be very intimately involved in the events occurring at the shoot apex during flower initiation. The flowering stimulus, formed in the leaves of photoperiodic plants, must, therefore, act as the shoot apex by affecting the gene-switching processes and thus involve nucleic acid metabolism.

Nucleic acid changes occurring at the shoot apex during flower initiation have been studied in two ways: (1) by following the incorporation of radioactive precursors into RNA, and (2) by the use of inhibitors of RNA synthesis.

In the LDP, *Lolium temulentum*, there is marked incorporation of radioactive precursors into RNA (thus indicating active RNA synthesis) at the shoot apices on the morning of the day following a single LD cycle, which is precisely the time at which the flowering stimulus is estimated to arrive at the apex. These changes in RNA and protein synthesis are most prominent in the cells on the flanks of the meristem which are destined to give rise to the spikelets (Fig. 10.6). Similarly, in the LDP, *Sinapis alba* (mustard), which will also flower in response to a single LD cycle, there is a marked increase in RNA synthesis in the central and peripheral zones of the apical meristem at about 17 hours following the beginning of the LD cycle. Later, active DNA synthesis occurs, and this is followed by mitosis.

The need for RNA and DNA synthesis has also been studied by using antimetabolites, such as 2-thiouracil (2TU), which inhibits RNA synthesis, and 5-fluorodeoxyuridine (5FDU) which inhibits DNA synthesis. Thus, application of 2TU to the shoot apex of *Sinapis alba* is most inhibitory to flowering when applied during the twelfth to twentieth hours after the beginning of a single LD cycle, suggesting that RNA synthesis during this period is an essential requirement for flower initiation at the shoot apex. Comparable results have been obtained with the SDP, *Xanthium* and *Pharbitis*. By studying the effects of 5FDU, it also appears that DNA synthesis is essential for flower initiation in these two latter species. However, in *Pharbitis*, 5FDU is effective in inhibiting flower initiation even if applied 24 hours after the end of an inductive dark period (i.e. after the arrival of the flower hormone), suggesting that it may inhibit the later formation of flower primordia, rather than the processes initiated by the flowering stimulus on its arrival at the shoot apex.

The results of these studies are fully consistent with the view that the initial events associated with the arrival of the flowering stimulus at the shoot apex involve marked changes in the nucleic acid metabolism, which might be expected if there is a "reprogramming" of gene activity during the change from vegetative to reproductive development.

NUTRITION AND FLOWERING

Farmers and gardeners have long known that fertilizers and manures have a marked effect on the balance between vegetative and reproductive growth. Indeed, for many practical purposes there seems to be an antagonism between vegetative growth and reproduction, and manurial treatment which favours strong vegetative growth may be unfavourable for flowering and fruiting. Experimental studies give some support for this view. Thus, it is found that low levels of nitrogen tend to result in earlier flowering in certain long-day plants. It has been shown that high nitrogen and carbohydrate nutrition delay flower initiation in pea (*Pisum sativum*). However, the number of cases in which a clear effect of mineral nutrition on the onset of flowering has been demonstrated is rather small. Mineral nutrition appears to have an important effect on flower initiation in fruit trees, high levels of nitrogen tending to promote vegetative growth and reduce flowering.

FLOWERING IN "NEUTRAL" SPECIES

In a large number of species the onset of flowering is a daylength or chilling response, and the discovery of photoperiodism and vernalization represents a very real advance in our understanding of the physiology of flowering. However, it must be remembered that in many other species, probably equally numerous, flowering is not greatly affected by daylength or winter-chilling (Table 10.2). This is the group referred to earlier (p. 199) in which flowering is relatively insensitive to external conditions, and although in these species the length of the vegetative phase may be modified by environmental conditions, the effects of the latter are not completely overriding. However, such species, which will be referred to as "neutral" species, are not sharply distinguished from those showing a "quantitative" response to daylength, which have been referred to as "facultative" LDP or SDP (p. 202).

TABLE 10.2. Some examples of species showing "neutral" flowering responses

<i>Cucumis sativus</i> (Cucumber)	<i>Phaseolus vulgaris</i> (Dwarf bean)
<i>Fagopyrum tataricum</i> (Buckwheat)	<i>Poa annua</i> (annual meadow grass)
<i>Fuchsia hybrida</i> (Fuchsia)	<i>Rosa</i> spp. (Rose)
<i>Helianthus annuus</i> (some varieties)	<i>Senecio vulgaris</i> (Groundsel)
<i>Lathyrus odoratus</i> (Sweet pea)	<i>Solanum tuberosum</i> (Potato)
<i>Lycopersicon esculentum</i> (Tomato)	<i>Vicia faba</i> (Broad bean)
<i>Nicotiana tabacum</i> (Tobacco, certain varieties)	

In these neutral species, flowering appears to be determined primarily by some *internal* mechanism, since even when grown under constant environmental conditions a species such as sunflower will grow vegetatively for a certain period and then become reproductive; thus the transition cannot be caused by any change in external conditions and must be

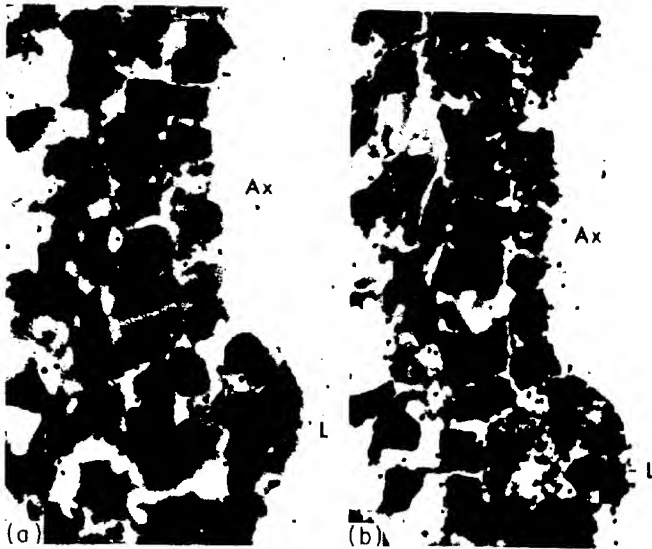


FIG. 10.6. Autoradiographs of vegetative and induced shoot apices of *Lolium temulentum*, labelled with ^3H -otic acid. A. Axillary bud site (Ax) and leaf primordium (L) in a vegetative (short-day) apex. B. Axillary bud site of plant which has been exposed to long day, in which there has been active incorporation of otic acid (as shown by the density of silver grains), indicating active nucleic acid metabolism. (From R. B. Knox and I. T. Evans, *Austr. J. Biol. Sci.* **21**, 1083, 1968.)

regulated by some "internal" mechanism. Moreover, it would appear that the transition from the vegetative to the flowering condition in such species is but one manifestation of a more general phenomenon, since progressive changes during development are very common, as shown by the development of morphological differences in successive organs such as leaves. It is commonly found that there are changes in the size and shape of successive leaves, the seedling or primary leaves being smaller than the later ones, and in species with deeply indented or compound mature leaves the primary leaves are usually much simpler in form (Fig. 10.7) with a progressive series of later formed leaves showing increasing segmentation. Plants showing such changes are said to exhibit *heteroblastic development*. These changes may be affected by environmental factors, such as light intensity and mineral nutrition, but they are not *dependent* on environmental changes and will occur even under constant conditions.

The changes in leaf shape appear to reflect progressive changes in the size and shape of the shoot apex. For example, in the sunflower plant the diameter of the sub-apical region increases progressively as the plant grows and ultimately these changes are terminated by

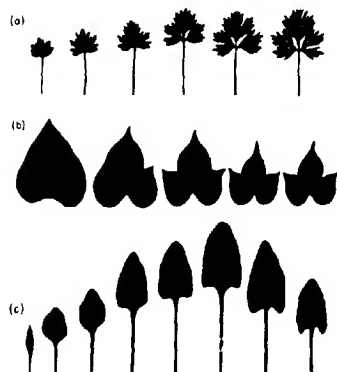


FIG. 10.7. Heteroblastic development, as shown by successive leaves of (a) *Delphinium ajacis*, (b) Morning glory (*Ipomoea hederacea*), (c) sugar beet (*Beta vulgaris*). (From E. Ashby, *New Phytologist*, **47**, 153, 1938.)

the formation of an inflorescence. It seems likely that the changes in leaf shape result from these changes in the shoot apex, leaf primordia produced on a larger apex apparently being capable of continuing their development for a longer period and hence producing a more mature leaf form.

This heteroblastic development appears to be an indication of the progress of the plant towards maturity and the attainment of the reproductive condition. For example, early flowering varieties of the cotton plant (*Gossypium*) show a steep gradient in leaf shape changes, whereas in later-flowering types the rate of change in leaf shape is less steep.

Evidence of progressive changes towards the flowering condition has come from experiments in which internode segments of tobacco were taken and grown in sterile culture. These segments produced callus and regenerated buds. Internode segments from young plants or from the lower region of the stem of older plants produced vegetative buds, whereas stem segments from the upper part of a flowering plant produced flower buds, with few leaves or bracts. Segments taken from an intermediate region of the stem first produced leaves and bracts and then a flower. Thus, there appeared to be a gradient in the propensity to produce flowers, from the lower to the upper part of the plant.

The nature and causes of the changes occurring in the shoot apex during development are completely unknown but there would appear to be at least three general possibilities:

- (1) There is some inherent pattern of behaviour of the apex, which will pursue its appointed course, independently of the differentiated portions of the plant.
- (2) The behaviour of the apex is affected and determined by influences arising in the mature parts of the plant, e.g., it might depend upon the attainment of a certain minimum leaf area.
- (3) The behaviour of the apex may be dependent upon the gradual accumulation of

certain metabolites which must attain a certain threshold concentration before flowering will occur.

At present, there is no incontrovertible evidence to enable us to decide which of these hypotheses is correct. Attempts have been made to obtain evidence on this problem in experiments with sunflower, in which seedlings tips were grafted on to stocks of various ages and tips from older plants were grafted on to seedlings. In the first type of graft it was found that seedling tips formed fewer nodes than they would have done on their parent plants. On the other hand, grafts of tips from older plants on to seedlings formed a greater number of nodes than they would have on the parent plants. These results would seem to indicate that the behaviour of the apex is determined by influences from the mature parts of the plant.

It will be remembered that among plants which are sensitive to daylength or chilling, there are some species which will not respond until they have reached a certain minimum size ("ripeness-to-flower"), and before they have reached this stage seedling plants may be described as in a "juvenile phase". This phenomenon would seem to correspond to the necessity for neutral species to reach a certain size before flowering. Indeed, photoperiodic and vernalizable species will behave like neutral species if they are maintained from germination under constant conditions favourable to flowering, in that they will undergo a certain period of vegetative growth and then commence to initiate flowers. Thus, it would appear that the physiology of flowering is not qualitatively different in the "sensitive" and "neutral" groups of plant, but rather that they differ in *degree* of sensitiveness to external conditions.

If we are correct in assuming that the difference between the flowering responses of "sensitive" and "neutral" species is primarily one of degree, then it might be assumed that a flower hormone is involved in neutral as well as in sensitive species, but there is little evidence as yet for the occurrence of flower hormones in neutral species.

THE DIVERSITY OF FLOWER-CONTROLLING FACTORS

As we have now seen, flowering is affected and controlled, in various species, by a range of factors, some of which are external to the plant, such as daylength, temperature and nutrition, and others which arise within the plant itself, as seen in day-neutral species. There is a bewildering mass of facts on the effects of these various factors and it is difficult to see how these parts of the "jigsaw puzzle" fit together to give an overall, unified picture. It might be asked whether the fact that quite different environmental factors may control flowering implies a different control mechanism for each factor. In attempting to answer this question it is useful to approach the problem from a different viewpoint and to ask not, what makes a plant flower under a given set of conditions, but rather, why they *fail* to flower under a different set of conditions. Thus, flowering may be prevented by a number of different factors in various species, although the flower-promoting processes might be the same in these species.

For example, in apple trees, flower initiation is rather sensitive to mineral nutrition, but not to photoperiod. We do not need to postulate that flower hormone synthesis depends upon low nitrogen levels—it may well be the case that the “flower hormone” is always present in apple shoots, but that under conditions of high nitrogen nutrition there are high levels of, for example, endogenous gibberellins, which are known to inhibit flowering in this plant. On the other hand, in *Xanthium* it would appear that synthesis of the flower hormone is blocked under long days. In plants requiring vernalization, yet another step in the flowering processes may be blocked in unchilled plants. From this viewpoint, under favourable conditions the flowering processes may be blocked in different ways and at different steps in plants of various response types. Similarly, as we have seen, flowering may be blocked by different factors in the same plant at different stages of its life cycle. In the seedling stages flowering may be prevented because the plant is still in a juvenile phase and is not yet capable of responding to favourable environmental conditions. But even when it has attained ripeness to flower, it may be prevented from flowering by unfavourable environmental conditions.

It is not difficult to envisage that in some cases flowering does not occur because the stimulus is not being synthesized in the leaves, while in other cases the limiting step may be the inability of the shoot apices to respond to the stimulus, possibly because of inappropriate levels of gibberellins.

FLOWERING IN WOODY PLANTS

So far, our consideration of the physiology of flowering has been restricted to herbaceous plants. Flowering in woody plants presents a number of characteristic features which will be briefly described.

The first important feature to note is that there is considerable variation between species in the length of time elapsing between flower initiation and complete development of the flower. In some species, such as sweet chestnut and many other late-summer-flowering trees and shrubs, such as *Buddleia*, *Fuchsia*, *Hypericum*, *Hibiscus syriacus*, *Caryopteris*, etc., the flowers are formed on the current year's shoots and flower initiation is followed immediately by the further full development of the flower, as in herbaceous species, i.e. there is no gap between the early and later stages of flower development. In many other woody plants, however, especially temperate species, flower initiation occurs during the summer within resting buds formed earlier in the same year, but the development of the flower parts becomes arrested at an early stage and the further development and emergence of the flower does not occur until the following spring. This is the situation in a large number of common European woody plants, e.g. oak (*Quercus*), ash (*Fraxinus*), sycamore (*Acer*), elm (*Ulmus*), pine (*Pinus*), apple, plum, peach, black currants, gooseberries, etc. In such species the buds containing flower primordia become dormant in the late summer or autumn and require a period of exposure to chilling temperatures to break the dormancy (p. 259). Once the dormancy has been broken by winter chilling they become capable of emerging as the

temperatures rise in the spring. In some winter- or early spring-flowering trees and shrubs the buds containing flowers are able to grow at lower temperatures than the vegetative buds, so that the flowers may actually emerge before the leaves, e.g. hazel (*Corylus*), willow (*Salix*), jasmine (*Jasminum nudiflorum*), almond (*Prunus persica*), elm, ash, oak, etc.

We have very little information regarding the effect of environmental factors on flower initiation in woody plants. This is mainly due to the technical difficulties in experimenting with mature trees. It is not possible to work with seedling trees since these are still in the juvenile phase and are not capable of flowering (p. 250). However, it is possible to study the effects of environmental conditions on flowering of trees by taking scions from mature trees and grafting them on to seedling stocks. In this way it is possible to obtain small trees which are potentially capable of flowering. In a few woody plants, flower initiation appears to be controlled by daylength conditions. Thus, long days are necessary for flower initiation in birch (*Betula*), *Erica* and *Calluna*. On the other hand, short days promote flower initiation in coffee (*Coffea arabica*), black currant (*Ribes nigrum*), *Poinsettia* and *Hibiscus*. Flower initiation in other woody plants, e.g., pine, larch, beech, apples, cherries, plums, appears to be unaffected by photoperiodic conditions, which, nevertheless, have a profound effect on vegetative growth of some of these species (p. 257).

In general, vernalization does not appear to play an important role in flower initiation in woody plants, with a possible exception of the olive tree (*Olea europaea*), which apparently initiates flowers in response to cool temperatures in the winter.

On the other hand, there is no doubt that temperature, rainfall and soil nutrients may have a profound effect on the flowering of many woody plants. It is well known that a hot, sunny summer is frequently followed by abundant flowering of many tree species in the following spring, and the flowering of beech has been shown to be particularly affected in this way. Evidently high temperatures and possibly high light intensity (and hence the formation of abundant carbohydrate reserves) favour flower initiation in many tree species. The effect of mineral nutrition on flower initiation in fruit trees was mentioned earlier (p. 244).

A number of instances have been reported in which flower initiation in woody plants has been influenced by gibberellins and by synthetic growth retardants. Thus, "flowering" has been stimulated by gibberellic acid in a number of conifers, including *Cupressus*, *Chamaecyparis*, *Juniperus* and *Thuja*. By contrast, gibberellic acid inhibits flowering in apple, pear, black currant, grape (*Vitis cinifera*), *Syringa*, *Fuchsia*, and a number of other woody plants. On the other hand, growth retardants such as "CCC" (chlorocholine chloride) and "Phosfon D", which apparently inhibit gibberellin biosynthesis in plant tissues promote flowering in apples, pears and azaleas (*Rhododendron* spp.).

PHASE CHANGE IN WOODY PLANTS

We have seen that in "day-neutral" annual species the onset of flowering is apparently regulated by some endogenous mechanism, the nature of which is unknown, but that the

effect of this regulatory mechanism is that the plant does not flower until it has attained a certain size. This "size effect" is seen also in species showing daylength or chilling responses, so that there is a juvenile phase during which flowering cannot be induced. An apparently analogous phenomenon is seen in woody plants, the seedlings of which normally show a juvenile phase during which they make active growth but remain vegetative. The transition to the flowering condition occurs only after a delay which varies greatly from species to species, ranging from 1 year in certain shrubs to 30–40 years in forest trees such as beech (Table 10.3). Once a given tree commences flowering, it normally continues to do so every year, although, as we have already seen (p. 249), in species such as beech, flowering is sensitive to weather conditions and may occur irregularly. Thus, on the basis of the flowering behaviour we may distinguish between a *juvenile* and an *adult* (or *mature*) phase in the life history of the tree.

TABLE 10.3. Duration of juvenile period in forest trees

	years
<i>Pinus sylvestris</i> (Scots pine)	5–10
<i>Larix decidua</i> (European larch)	10–15
<i>Pseudotsuga taxifolia</i> (Douglas fir)	15–20
<i>Picea abies</i> (Norway spruce)	20–25
<i>Abies alba</i> (Silver fir)	25–30
<i>Betula pubescens</i> (Birch)	5–10
<i>Fraxinus excelsior</i> (Ash)	15–20
<i>Acer pseudoplatanus</i> (Sycamore, Maple)	15–20
<i>Quercus robur</i> (English oak)	25–30
<i>Fagus sylvatica</i> (Beech)	30–40

Differences between juvenile and adult stages are seen not only in flowering behaviour but also in various vegetative characters. Thus, in certain species, such as ivy (*Hedera helix*), mulberry (*Morus*), *Acacia*, *Eucalyptus* and juniper, the leaf shape in the juvenile phase is very different from that of the adult stage. In oak and beech there is a marked tendency for the dead leaves to be retained on the shoots of juvenile trees during the winter, whereas in the adult stages they are shed normally. In some species the juvenile and adult stages are distinguished by marked differences in phyllotaxis, as in ivy (see below). Another morphological character which changes during ontogeny is the development of thorns—for example, in lemon trees the juvenile stages are commonly more thorny than the adult.

Among the physiological differences observed between juvenile and adult stages is the rooting ability of cuttings: it is very commonly found that whereas cuttings from young trees root readily, after the parent trees attains a certain age this rooting ability is greatly diminished or entirely lost.

An interesting feature of these phase differences is that the lower parts of the tree retain the juvenile stage after the upper parts have developed adult characters. This can be seen very well in ivy, where the lower regions of an erect-growing vine show the palmate type of leaf and are purely vegetative, whereas the upper parts show ovate leaves and normally produce abundant flowering shoots.

Not only is there apparent stability of the juvenile and adult phases in the same plant body, but cuttings taken from different regions and rooted retain their juvenile or adult characters for a long period. This is well known in ivy, for example, where cuttings taken from the juvenile part of the vine had palmate leaves with opposite phyllotaxis, the shoots are trailing and are coloured with anthocyanin and produce abundant adventitious roots, but do not flower; cuttings of adult shoots, on the other hand, produce plants with ovate leaves, spiral phyllotaxis, and the shoots are erect, green and produce few, if any, adventitious roots, but flower readily (Fig. 10.8). Cuttings from adult ivy shoots may grow for many years and produce shrubs known to gardeners as "tree ivy".



FIG. 10.8. Cuttings taken from adult part of parent vine of ivy (*Hedera helix*). These cuttings will continue to retain for several years the "adult" characters, such as leaf shape, phyllotaxis, and the capacity to flower, although high temperatures favour reversion to the juvenile condition. (Print supplied by Dr. L. W. Robinson.)

A similar retention of juvenile and adult characters is seen in grafting experiments. Thus, scions from flowering regions of mature trees continue to flower when grafted on to quite small seedling stocks. This is a common observation in forest tree breeding and is known for species such as birch, larch and pine. Likewise, scions from mature fruit trees grafted on to suitable root stocks flower readily, whereas scions from young seedlings treated in the same way show delayed flowering.

The phenomenon of phase change, involving stable, non-genetic changes which can be transmitted through many cell-generations, shows certain interesting parallels with the changes occurring during vernalization. This subject will be discussed further in Chapter 13.

FURTHER READING

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CHAPTER 11

Dormancy

OUTSIDE the equatorial regions there are seasonal variations in climatic conditions, which are most notable in the temperate zones. These variations are especially marked with respect to light intensity, daylength, temperature and frequently also to rainfall. As a result there is a regular alternation of seasons favourable and unfavourable for growth and this alternation has had a marked effect on the pattern of the life cycles evolved by the higher plants. The necessity to withstand low temperatures during the winter and, in some regions, hot dry conditions during the summer, poses special problems for the plant, and we now have to consider some of the ways in which these problems have been met.

THE BIOLOGICAL SIGNIFICANCE OF DORMANCY

Plant cells normally contain a large amount of water which is liable to freeze at low temperatures, with grave risk of damage to the protoplasm. Tropical plants are very easily killed by freezing, but it is evident that plants of temperate and arctic regions must have become adapted to survive the period of winter frost—they have developed *cold-resistance*. Although cold-resistance has been studied for many years, our understanding of its biochemical basis is still far from complete, and a discussion of this subject here would take us too far afield.

In many cold-resistant species the general morphological appearance of the plant during winter is not essentially different from that in the summer—the growth rate of the plant is reduced or arrested during the winter, but the growing points of the shoots are still in a potentially active condition, and may make some growth during mild periods, as in many biennial plants. In such species the whole plant, including the apical meristems, is relatively cold-resistant. Other species, of course, show distinct differences between their summer and winter states. Thus, in woody plants the shoot apices cease active growth and become enclosed in bud scales, to form winter-resting buds. They are then said to have become *dormant*. Many woody plants are much more cold-resistant in the dormant than in the actively growing condition. Thus, seedlings of forest trees, such as larch (*Larix*) and *Robinia*, which continue growing late into the autumn, are very liable to be damaged by early

frosts, but if they have ceased growth and their growing points have entered the dormant condition, they then remain cold-resistant throughout the winter.

The reason why dormant buds should be more cold-resistant than actively growing tissues is not fully understood. However, it is fairly clear that the cold-resistance of dormant tissue is due to certain protoplasmic characters, and that it is not primarily due to the presence of the bud scales, the protective function of which is probably concerned with the reduction of water loss—one of the secondary effects of winter cold is to increase the difficulties of the plant in maintaining an adequate water balance. Under frosty conditions, especially when accompanied by wind, the plant continues to lose water, but is unable to take up replacement supplies if the soil is frozen. There is thus a considerable danger of damage from drought under winter conditions, but water loss is reduced by the enclosure of the growing parts within a covering of bud scales, and also, in deciduous trees, by the falling of leaves in autumn which reduces the total surface area over which evaporation can occur.

The danger of winter drought and low temperature seems to have influenced not only the evolution of woody plants, but has apparently had a profound effect on the form of many other types of plant. Many plants over-winter entirely below ground, as bulbs, corms and rhizomes; although such organs will be partly insulated against frost, they will also be protected against drying when the soil does become frozen to some depth. Some dormant organs, such as bulbs, are probably adaptations to hot, dry summer conditions, such as are found in the Mediterranean region.

Whereas perennial plants have developed special organs which resist the unfavourable conditions of winter, annual plants have pursued yet another course—they frequently over-winter in the form of seeds. The seeds of many annual plants, particularly of the common weeds of arable land, germinate almost immediately they are shed, if conditions of temperature and moisture are favourable. But the seed of many other plants does not germinate immediately (or only a proportion of the seeds do so) and remains in the soil until conditions become favourable for germination in the following spring. Now seeds are generally very much more cold-resistant than the growing plant of the same species. Dry seeds may resist freezing down to as low as -23°C . Some seeds, such as those of Leguminosae (clover (*Trifolium*), broom (*Cytisus*), *Laburnum*, etc.) do not, in fact, take up water immediately they are shed, due to the fact that they have a coat which is impermeable to water, and such seeds will be capable of withstanding very severe frost.

The majority of seeds imbibe water as soon as they fall on to moist soil, but, as already stated, they do not necessarily germinate immediately. Such imbibed seeds are less cold-resistant than in the dry state, but nevertheless many of them still retain a considerable degree of cold-resistance, and apparently certain annual species which are frost-tender in the actively growing state are able to survive the winter in the form of seed.

FORMS OF DORMANCY

Dormancy may be defined as a state in which growth is temporarily suspended. In some species the cessation of growth is directly due to unfavourable temperature and light con-

ditions; thus many pasture grasses remain in continuous growth throughout a mild winter and cease growth only when temperatures fall to about 0–5°C. Similarly, certain annual weeds, such as groundsel (*Senecio vulgaris*), chickweed (*Cerastium* spp.) and Shepherd's purse (*Capsella bursa-pastoris*), stop growing only during the coldest part of the winter. In such cases the dormancy of plants is evidently caused by the unfavourable external conditions, and in this case we speak of *imposed* or *enforced dormancy*.

However, in many cases the unfavourable conditions are not directly the cause of dormancy. Thus, many trees form winter-resting buds during the summer and autumn, when temperatures and light conditions are still favourable, and long in advance of the onset of winter. In such woody plants the cause of dormancy appears to lie within the tissues of the buds themselves, and we then speak of *innate* or *spontaneous dormancy*. This form of dormancy also occurs in many seeds. Thus, if freshly harvested barley grains are planted under warm, moist conditions, a high percentage of them will fail to germinate. If, however, the barley is stored dry for a few months, the seed will then be found to germinate readily when planted under the same conditions as previously. Thus, the failure of freshly harvested barley grains to germinate is not due to external conditions being unfavourable for growth, but must be due to some cause within the seed itself.

Innate or spontaneous dormancy is found not only in buds and seeds, but in other types of resting organs such as rhizomes, corms and tubers.

BUD DORMANCY IN WOODY PLANTS

The majority of temperate woody plants, including both coniferous and dicotyledonous species, show a well-marked dormancy or resting phase during the annual growth cycle and this is usually accompanied by the development of resting buds. The typical resting bud involves the "telescoping" of the bud scales and leaf primordia in the apical region, due to the arrest of normal internode extension. In some genera (e.g., *Betula*, *Fagus*, *Quercus*) having stipules, this telescoping of the shoot apical region leads to the formation of a resting bud, since the overlapping stipules in this region form the bud scales. In other species the protective scales represent leaves, which may be only slightly modified, as in *Viburnum* spp., or more highly modified, so that they frequently represent only the leaf base, as in *Acer*, *Fraxinus*, *Malus* and *Ribes* (Fig. 11.1). During the development of such buds, certain leaf primordia show greater marginal growth than occurs during normal leaf development, whereas lamina development is suppressed, and these primordia give rise to the bud scales. The younger leaf primordia, formed within the bud scales, are arrested at an early stage in their development, and give rise to normal leaves when the buds resume growth in the following spring. In some species, such as pines, growth of a bud may continue for several months from June to September. In some trees a terminal bud is not formed, since growth of the shoot is terminated by the death and abscission of the apical region, and growth is later continued from the uppermost axillary bud. Such species (which include *Tilia*, *Ulmus*,

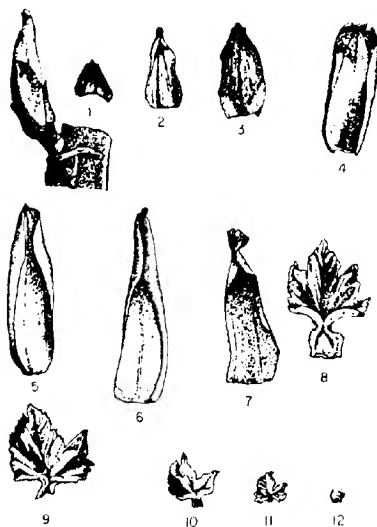


FIG. 11.1. Bud of *Ribes* and bud scales and leaves from a dissected bud. (From J. H. Priestley and L. I. Scott. *An Introduction to Botany*, 3rd ed., Longmans, Green & Co., London, 1955.)

Castanea, *Robinia* and *Ailanthus*) are said to show a *sympodial* (as opposed to a monopodial) growth habit.

When terminal buds are first formed they can frequently be induced to resume growth by various treatments including defoliation, either by hand or by insect attack. It would appear, therefore, that at this stage the terminal buds are not themselves innately dormant, but their growth is apparently inhibited by the mature leaves on the shoot. Similarly, lateral buds may be inhibited by the leaves, or by the main apical region in actively growing shoots, and are thus held in check by correlative inhibition (p. 130) rather than by innate dormancy. This phase of bud development is referred to as *summer dormancy* or *predormancy*. Later, in many species, the buds are found to have entered a state referred to as *true dormancy*, *winter dormancy* or *rest*. When they have entered this condition the buds will no longer resume growth if the shoots are defoliated, so that they are now innately dormant and not simply held in check by environmental conditions or inhibitory influences within the plant itself, as is the case during predormancy.

After a certain period of true dormancy the buds become capable, in the later part of the winter or early spring, of resuming growth when external conditions, particularly temperature, are favourable for growth. Thus, at this stage the buds are no longer innately dormant, but nevertheless for some time they may fail to grow because of low outdoor temperatures. This phase is referred to as *post-dormancy*. We shall now consider some of the

environmental factors controlling the development and breaking of dormancy in temperate woody plants, of which daylength and temperature are the most important.

The Development of Bud Dormancy

One of the most important factors affecting and controlling the induction of dormancy in woody plants is daylength. In the majority of species so far studied, long days promote vegetative growth and short days bring about the cessation of extension growth and the formation of resting buds in seedlings of woody plants (Fig. 11.2). However, a number of



FIG. 11.2. Photoperiodic control of bud dormancy in seedlings of birch (*Betula pubescens*). Plants transferred to short days (left) have ceased growth and formed resting buds, whereas seedlings maintained under long days will continue to grow actively for many months.

common cultivated fruit trees (*Pyrus*, *Malus*, *Prunus*) and certain other species, including the family Oleaceae, appear to be relatively insensitive to daylength changes.

The seedlings of some species, e.g. black locust (*Robinia pseudacacia*), birch (*Betula pubescens*) and larch (*Larix decidua*), can be maintained in continuous growth for at least 18 months under LD conditions in a warm greenhouse, whereas under SD they cease growth within 10–14 days. On the other hand, other species, such as sycamore (*Acer pseudo-platanus*), horse chestnut (*Aesculus hippocastanum*) and sweet gum (*Liquidambar styraciflua*), show delayed dormancy under LD, but they cannot be maintained in growth indefinitely under these conditions. For those species which can be maintained in continuous growth under LD, there appears to be a certain critical daylength below which dormancy is induced and above which dormancy does not occur.

As in the flowering responses of herbaceous plants, the photoperiodic responses of woody seedlings appear to depend upon the length of the dark period, rather than of the photoperiod, and if a long dark period is interrupted by a short "light-break", the effect of the dark period is nullified and dormancy is delayed. The most effective region of the spectrum for this light-break effect lies in the red, suggesting that phytochrome is involved, and clear red/far-red reversibility has been demonstrated for seedlings of larch.

The response of woody seedlings depends on the daylength conditions to which the leaves are exposed. In sycamore, as in herbaceous species, it is the young, fully expanded leaves which are the most sensitive to daylength, but in birch seedlings even quite young leaves in the apical regions show sensitivity to photoperiod.

How important are these photoperiodic responses in determining the formation of resting buds and the onset of dormancy in nature? It has been shown that the seasonal decline in daylength is important in determining the onset of dormancy in seedlings of species which normally continue active growth into the autumn, e.g. *Larix decidua*, *Populus* spp. and *Robinia pseudacacia*. But it is very common to find that older trees show a very much shorter period of extension growth than do seedlings of the same species, and they frequently cease growth in June or July, when the natural photoperiods are still long. In such cases, it seems doubtful whether declining daylength is important in determining the formation of resting buds, and it seems more likely that some change, in either nutrient levels or in hormonal balance, arising within the tree itself, determines the period of growth and the onset of dormancy. However, as we have seen, resting buds are at first in a state of predormancy and only later enter a state of true dormancy; it is possible that the declining daylength in the autumn plays a role in the transition of the buds from predormancy to true dormancy.

It has been found that leaf fall in some woody plants is promoted by short days, and delayed leaf fall is sometimes observed in trees growing near street lights. However, in nature, low temperatures and possibly low light intensity are probably at least as important as daylength in determining the onset of leaf senescence and abscission.

In addition to the observable morphological changes associated with the induction of dormancy by short days there are also biochemical changes which are reflected in increased cold-resistance. Seedlings of black currant (*Ribes nigrum*) and *Robinia pseudacacia* which

have been exposed to SD are markedly more cold-resistant than seedlings which have been grown throughout under LD. Cold acclimatization in *Cornus stolonifer* depends upon exposure to both SD and decreasing temperature.

It is now well established that wide-ranging woody species, such as *Pinus sylvestris* and *Picea abies*, show marked ecotypic differences in photoperiodic responses in relation to both the latitude and the altitude at which they occur naturally. Northern races are frequently found to require longer photoperiods for active extension growth than do more southern races, adapted to shorter natural photoperiods. This fact suggests that woody plants are rather closely adapted to natural daylength conditions, and that the latter probably play an important controlling role in the seasonal cycle of growth and dormancy.

Emergence of Buds from Dormancy

Usually the terminal resting buds formed during the summer or fall remain dormant until the following spring, when they expand and form new shoots. The dormancy of the buds diminishes during the course of the winter, as can be demonstrated very simply by collecting twigs of trees such as lime (*Tilia*), sycamore (*Acer pseudoplatanus*), poplar (*Populus*) and willow (*Salix*) at different times during the winter and placing them in water in a warm room or greenhouse. It is found that twigs collected in October, November and early December usually remain dormant when they are brought into warm conditions. A fairly high proportion of buds collected in January will be found to expand in 2-3 weeks, and with later dates of collection, e.g. February or March, the buds burst increasingly rapidly after they are brought into the warm.

Many woody plants require to be exposed to a period of winter chilling to overcome the dormancy of their buds, as can be shown by growing small trees of species such as poplar or sycamore in pots and, when they have become dormant in the autumn, keeping some of them out-of-doors throughout the winter and some of them in a warm room or greenhouse. In the spring, the buds of young trees which have been kept outdoors will expand in the normal way, but those which have been kept in the warm will still be dormant and may remain so until well into the summer; indeed, a certain proportion may ultimately die without ever resuming growth. Temperatures in the range 0-5°C are the most effective in overcoming bud dormancy, and chilling periods of 260 to 1000 hours are required. In regions with cold winters, the chilling requirements are normally fully met by the spring, but in warm climates, such as those of California and South Africa, where the winters are very mild, difficulties may be met in cultivating certain fruit trees, such as peaches (*Prunus persica*), since the chilling requirements of the buds may not be met and delayed and irregular bud-break may occur in the spring.

It should be noted that although chilling is necessary to remove the dormancy of the buds of many trees, warm temperatures are necessary for the growth of the buds after chilling. Frequently the chilling requirements are met by January, but in many regions

the buds may fail to resume growth then because the temperatures are still too low, and they remain in the phase of postdormancy. Thus, the time of bud burst in the spring is normally determined by the return of warmer conditions.

In the majority of North Temperate woody species so far studied, once bud dormancy has been fully induced by SD, they cannot be induced to resume growth by transfer to LD, and the dormancy can normally be overcome only by chilling. However, in a few species the unchilled buds can be induced to resume growth under LD or continuous illumination. Thus, if leafless seedlings of beech (*Fagus sylvatica*), birch (*Betula* spp.) or larch (*Larix decidua*) are placed under continuous light in a warm greenhouse in the autumn, the buds will soon expand.

At first sight it would seem that the response of dormant buds to photoperiod contradicts the rule that it is the leaves which are the organs of photoperiodic "perception", and that the apical meristematic region is insensitive to daylength. However, it should be remembered that resting buds contain well-developed leaf primordia and that therefore the differences between species, such as beech, and other species relates primarily to a difference in the age at which the leaves become sensitive to photoperiod.

It is not clear whether the photoperiodic control of bud-break is important in nature, but there is evidence that bud-break in *Fagus sylvatica* may be dependent upon lengthening daylengths in the spring, although in many regions temperature is also likely to be a limiting factor for this species. In *Rhododendron*, also, bud-break appears to be determined by daylength.

DORMANCY IN VARIOUS ORGANS

Various other types of organ, such as rhizomes, corms, bulbs, tubers and the winter-resting buds of aquatic plants, show dormancy. In the aquatic plants *Stratiotes*, *Hydrocharis* and *Utricularia* the dormancy of the winter-resting buds is induced by short days, in association with high temperature. Short days also promote the formation of resting buds in the insectivorous plant, *Pinguicula grandiflora*, and the dormancy of the buds is overcome by chilling.

By contrast, dormancy in bulbs of onion (*Allium cepa*) is promoted by LD so that the bulbs develop and "ripen" in the summer. The period of dormancy of onion bulbs is shortest when the bulbs are stored at cool temperatures.

The rhizomes of lily-of-the-valley (*Convallaria majalis*) normally become dormant during the summer and they require a 1-week period of chilling at 0.5–2°C, or 3 weeks at 5°C to remove this dormancy. Similarly, when *Gladiolus* corms are exposed to warm soil temperatures they do not grow, but periods as short as 24 hours at 0–5°C will break their dormancy.

Although the tubers of some varieties of cultivated potato have a chilling requirement for dormancy breaking, this is apparently not the case with all varieties.

ARTIFICIAL MEANS OF BREAKING BUD DORMANCY

A very wide range of treatments, especially using various chemicals, has been found to break the dormancy of resting organs. One of the simplest methods of breaking the dormancy of woody plants consists of immersing the shoots in warm water (at 30–35°C) for 9–12 hours; this treatment is used by florists for “forcing” the flower buds of lilac and *Forsythia*, to get blooms much earlier than normally. Exposure to ether vapour is also effective in removing the dormancy of lilac buds and of lily-of-the-valley rhizomes, the flowers of which can thus be obtained very early in the winter for sale by florists. Among the other substances which have been found very effective in breaking dormancy are thiourea and ethylenechlorhydrin, which will remove the bud dormancy of a wide range of woody plants and are also effective with potato tubers and rhubarb root stocks.

As will be shown below, various growth regulators, including gibberellins, cytokinins and ethylene, will also break the dormancy of buds and seeds of many species.

DORMANCY IN SEEDS

Morphologically, the seed consists of an embryo surrounded by one or more covering structures, of which the most important is the testa, which is usually derived from the integuments of the ovule. Some seeds contain a well-developed endosperm, which lies within the testa and may surround the embryo, or may lie to one side of it. Functionally, the seed is a “propagule” or dispersal unit, i.e. an organ of propagation. In many species the seeds are liberated from the fruit and the isolated seeds become the dispersal units. In other species, however, the fruits may contain a single seed which is retained within the fruit coat (pericarp), the fruit itself being shed as a whole and becoming the dispersal unit. Examples are provided by achenes, nuts, caryopses and so on, the precise definition of which does not concern us here. Although these latter structures are distinguishable morphologically from seeds, they perform the same biological function as seeds, i.e. they are functional propagules and hence it is convenient to refer to all such structures as seeds, although it is not strictly accurate to do so.

Although, as we shall see, dormancy in seeds shows many parallels with that in buds and other organs, the presence of enclosing coats introduces complications not found in buds, and we find several types of seed dormancy, which do not appear to correspond to any form of bud dormancy.

Hard Seed Coats

The seeds of certain families, such as the Leguminosae, Chenopodiaceae, Malvaceae and Geraniaceae, possess testas which are impermeable to water, so that such seeds are liable to lie dormant in the soil for considerable periods before germination occurs. Water uptake

by these seeds can be brought about by various treatments, such as abrasion by sand, treatment for short periods with concentrated sulphuric acid, etc., which remove the impermeable outer layer of the testa and permit penetration of water to the embryo. Seedsmen treat clover seed by rotating it in a drum lined with carborundum. Probably under natural conditions the activities of micro-organisms in the soil slowly break down the outer layers of the testa and so render water uptake possible.

Immaturity of the Embryo

In certain seeds the embryo is still immature when the seed is shed and germination cannot occur in such seeds until the embryo has undergone development. This is true of the seeds of wood anemone (*Anemone nemorosa*), lesser celandine (*Ficaria verna*), marsh marigold (*Caltha palustris*), ash (*Fraxinus excelsior*), and other species. In order for this further development to take place, the seeds must be imbibed with water and maintained under favourable temperature conditions. The time required for the embryo to complete its development may vary from about 10 days in *C. palustris* to several months in *F. excelsior*.

After-ripening in Dry Storage

The seeds of many species fail to germinate if sown immediately after harvesting, even though the embryo is fully mature. If they are stored dry at ordinary room temperatures, however, they gradually lose their dormancy and become capable of germinating when provided with suitable conditions (Fig. 11.3). This effect is called "after-ripening in dry-storage", and is found in several types of cereal, e.g. barley, wheat, oats and rice. The duration of the dormant period may range from a few weeks to several months. Other species

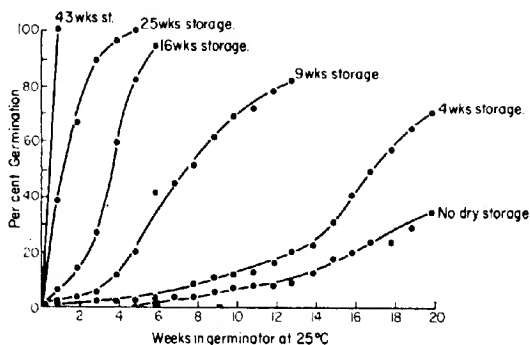


FIG. 11.3. Effect of after-ripening in dry storage at room temperature on germination rate of seeds of *Impatiens balsamina*. (From W. Crocker, *Growth of Plants*, Reinhold, New York, 1949.)

showing this type of dormancy include many grasses, black mustard (*Brassica nigra*), evening primrose (*Oenothera* spp.), clover (*Trifolium* spp.) and cultivated varieties of lettuce.

It is not known what causes this type of dormancy nor what are the changes which occur during the storage period which ultimately release the seed from dormancy. It would seem that the processes involved during this period of after-ripening are not of a metabolic nature, since they occur even in the dry seed, when metabolism is at a very low level.

This type of dormancy is of considerable economic importance in cereals. In regions where the weather during the harvest period is liable to be wet, dormancy of the grain is an advantage, since varieties which show such dormancy are less liable to germinate in the ear under wet conditions. From this point of view, dormancy is a desirable economic "character", which is deliberately selected for by plant breeders. On the other hand, for the production of malt from barley, dormancy of the grain is often a major problem, since it may be impossible to germinate the grain for several weeks after harvesting.

Light-sensitive Seeds

One of the most interesting forms of dormancy, and one which has received intensive study in recent years, is that shown by light-sensitive seeds. In a considerable number of species, exposure to light is necessary for germination, e.g. seeds of tobacco (*Nicotiana* spp.), foxglove (*Digitalis purpurea*), hairy willow-herb (*Epilobium hirsutum*), purple loosestrife (*Lythrum salicaria*), dock (*Rumex crispus*), and many others. On the other hand, in certain other species germination is inhibited by light, although the number of such species is considerably smaller than for light-promoted seeds; among the known light-inhibited seeds are those of love-in-a-mist (*Nigella*), *Nemophila*, *Phacelia* and annual phlox (*Phlox Drummondii*).

Light-sensitive seeds will only respond to light after they have imbibed water. The duration of illumination required by light-promoted seeds is often very short; for example, a high percentage of germination is obtained with lettuce seed exposed to only 1–2 minutes of light, while with seed of purple loosestrife a light-flash of only 0.1 seconds duration has a marked effect in stimulating germination. The responses of light-sensitive seeds are strongly affected by temperature and many seeds which are light-requiring at, say, 25°C become capable of germinating in the dark at lower temperatures, e.g., certain light-requiring varieties of lettuce. If certain light-requiring seeds are exposed to daily alternations of temperature, e.g., between 15°C and 25°C, they can be induced to germinate without exposure to light. Other treatments which can replace the light requirement of seeds include treatment with certain inorganic ions, especially nitrate, and certain organic substances, e.g., thiourea.

Many seeds which are light-requiring when freshly harvested gradually lose their light-requirement during storage and ultimately give full germination in complete darkness, e.g., light-sensitive varieties of lettuce. It would seem, therefore, that the changes occurring during after-ripening in dry storage in some way remove the light requirement.

As we have already seen, studies on the responses of light-sensitive varieties of lettuce seed played a key role in the discovery of phytochrome (Chapter 8, p. 183). It was shown that the red region of the spectrum promotes and far-red inhibits the germination of lettuce seed (Fig. 11.4). If red and far-red are given alternately, then whether germination occurs or not depends upon the nature of the last radiation to which the seeds were exposed. Thus, when phytochrome is converted to the P_{tr} form by red light, it evidently initiates a chain of processes which ultimately result in germination. It has been found that similar red/far-red responses are shown by other species of light-sensitive seeds and it seems probable that phytochrome is universally involved in light-promoted seeds.

Light-inhibited seeds have been very much less intensively studied than light-promoted ones, but it now seems probable that the same phytochrome system is involved in both types of seed, and that in the light-inhibited species the effect of far-red is enhanced so that it predominates over the effect of red. Thus, it has been shown that the light-inhibition of *Nemophila* seed is due mainly to the far-red region of the spectrum; red light, on the other hand, seems to have little or no promotive effect on this seed.

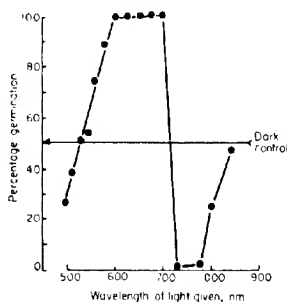


FIG. 11.4. Effect of red and far-red radiation on germination of lettuce seed. (From L. H. Flint and E. D. McAlister, *Smithsonian Inst. Misc. Collections*, **96**, 1-8, 1937.)

Dormancy Removed by Chilling

Gardeners have long known that the seeds of many species will not germinate if sown under warm conditions, but will lie dormant in the soil for long periods; if, however, they are sown out-of-doors in the autumn and exposed to winter conditions they will germinate in the following spring.

This behaviour led to the horticultural practice of "stratifying" the seed, i.e. placing it between layers of sand and leaving it out-of-doors during the winter. Such "stratified" seed is no longer dormant and germinates readily in the spring (Fig. 11.6). From such observations it is clear that exposure to winter cold is, in some way, necessary to break the dormancy of many seeds.

At one time it was believed that the dormancy of such seeds is due to hard and impermeable coats, and that freezing temperatures are necessary to break the coats. It is now known, however, that freezing temperatures are not required, and that, in fact, temperatures just above freezing (0–5°C) are more effective than lower temperatures (Fig. 11.5). Moreover, many seeds which have a chilling requirement do not, in fact, have hard coats, e.g., apple, birch.

The range of seeds showing a chilling requirement is very wide (Table 11.1) and includes both woody and herbaceous plants. In some seeds there is an obligate requirement for chilling, e.g., ash (*Fraxinus excelsior*), whereas in others, e.g., *Pinus* spp., a period of pretreatment at chilling temperature, although not essential, nevertheless increases and hastens subsequent germination. It should be noted that for chilling temperatures to be effective the seeds must be imbibed with water, there being no effect with dry seeds. The minimum period of chilling necessary to remove dormancy varies from species to species, but usually amounts to several weeks. In some species the embryo itself is dormant and can only be induced to germinate with difficulty if it is unchilled, e.g., mountain ash (*Sorbus aucuparia*), whereas in other species the embryo will germinate if the testa is removed and only the intact seed has a chilling requirement, e.g., sycamore (*Acer pseudoplatanus*). Seedlings grown from unchilled embryos frequently show "dwarfism", however, making sluggish growth and having very short internodes. This dwarfism of seedlings can itself be removed by chilling or by treatment with gibberellic acid.

Certain seeds, such as acorns (*Quercus*) and those of *Viburnum*, show "epicotyl dormancy"; such seeds germinate and develop a radicle in the autumn without any prior chilling but development of the epicotyl is dependent upon chilling, i.e. the epicotyl, but not the radicle, shows dormancy.

A few species have "2-year seeds", so called because they do not normally germinate until the second spring after shedding. Certain types of 2-year seeds have hard coats, as well as a chilling requirement, e.g., hawthorn (*Crataegus*) and *Cotoneaster*; because of the hard

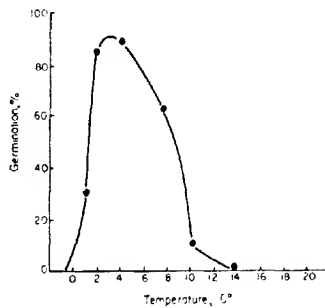


FIG. 11.5. Effect of chilling temperatures on dormancy of apple seed (germination after 85 days chilling at temperatures shown). (From P. G. de Haas and H. Scharder, *Zeitschrift für Pflanzenzüchtung*, 31, 457, 1952.)

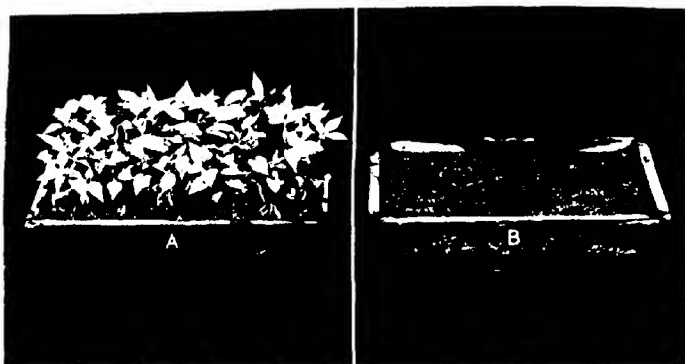


FIG. 11.6. Effect of winter chilling on dormancy of seeds of *Rhodotypos*. A. Seeds exposed to outdoor temperatures throughout the winter. B. Seeds maintained in a warm greenhouse throughout experiment. (Photograph supplied by late Dr. Lela V. Barton.)

TABLE 11.1. Woody species with seeds having a chilling requirement to overcome dormancy

<i>Acer</i> spp. (Maples)	<i>Malus</i> spp. (Apple and Crab apple)
<i>Betula</i> spp. (Birches)	<i>Picea</i> spp. (Spruce)
<i>Cornus florida</i> (Dogwood)	<i>Pinus</i> spp. (Pines)
<i>Corylus avellana</i> (Hazel)	<i>Prunus</i> spp. (including Peach)
<i>Crataegus</i> spp. (Hawthorns)	<i>Rosa</i> spp. (Roses)
<i>Fagus sylvatica</i> (Beech)	<i>Sequoiadendron giganteum</i> (Wellingtonia)
<i>Fraxinus</i> spp. (Ash)	<i>Tilia</i> spp. (Lime)
<i>Hamamelis virginiana</i> (Witch hazel)	<i>Thuja occidentalis</i> (Western red-cedar)
<i>Juglans nigra</i> (Walnut)	<i>Vitis</i> spp. (Grape)
<i>Liriodendron tulipifera</i> (Yellow poplar, Tulip tree)	

coats, the embryos are prevented from imbibing water as soon as shed, and hence chilling during the first winter is ineffective in removing dormancy. The hard coats are rendered permeable to water during the following summer, however, as a result of the activities of soil micro-organisms. When such imbibed seeds enter the second winter, the dormancy is broken and they become capable of germinating in the following spring.

The cause of the "2-year" behaviour is different in other species. Thus, seeds of lily-of-the-valley (*Convallaria*) and Solomon's seal (*Polygonatum*) require a chilling period to bring about growth of the radicle, but development of the epicotyl does not follow until the seeds have been subjected to a second winter's chilling.

The Role of the Coats in Seed Dormancy

The seed coats have been found to play an important role in the dormancy of the seed of many species. It has already been mentioned that although the embryos are dormant in

some seeds which have a chilling requirement, nevertheless in other species it is only the intact seeds which show dormancy, and the isolated embryos will germinate without chilling if the testa is removed. Similarly, certain light-requiring seeds, such as birch and lettuce, will germinate in darkness if the seed coverings are removed or even if they are only slit. Again, certain seeds which show a requirement for after-ripening in dry storage will germinate if the seed coverings are removed; for example, removal of the husks of barley, wheat, oats and rice will permit germination soon after harvesting, whereas when the coats are left intact such seeds normally require several weeks of after-ripening. It is clear, therefore, that the seed coats play an important role in at least three different types of dormancy, and, indeed, in all cases where the embryo itself is not dormant, the dormancy of the intact seed depends upon the presence of coats, which will include the testa, together with the endosperm and pericarp in some seeds. This conclusion raises the question as to the mechanism of these seed-coat effects.

It is possible that the seed coats present a physical barrier to gaseous exchange of oxygen and carbon dioxide between the embryo and the external air. It seems unlikely that seed coat effects are due to the accumulation of high internal concentrations of carbon dioxide since germination of lettuce seed is actually stimulated by keeping the seeds in high concentrations of this gas. On the other hand, several types of seed show higher oxygen requirements than do actively growing plants of the same species, suggesting that seed coats may present a physical barrier to oxygen uptake. The testas of marrow (*Cucurbita pepo*) seed have been shown to be much less permeable to oxygen than to carbon dioxide. Certain seeds can be induced to germinate either by slitting or removing the coats, or by maintaining the intact seed in a high concentration of oxygen, e.g., in *Betula* and non-after-ripened cereals. Studies on the respiration of germinating pea seeds suggest that anaerobic conditions may occur in the initial stages of germination until the testa is ruptured, when there is a marked increase in oxygen uptake (p. 278). Thus, several kinds of evidence support the view that seed coats may limit the uptake of oxygen.

Interference with oxygen uptake, especially in association with high temperatures, appears to be important in what is known as *secondary dormancy*. Thus, non-dormant seeds of *Xanthium* can be rendered dormant by embedding them in clay (which restricts gaseous exchange) and keeping them at 30°C for several weeks. Similar secondary dormancy phenomena have been shown for a number of other species, including members of the Polygonaceae and Rosaceae, e.g., apple and pear. In all these cases, the secondary dormancy can be overcome by chilling treatment, and it appears that the development of secondary dormancy is the reverse of after-ripening.

Secondary dormancy shows many resemblances to primary dormancy and it has been suggested by Vegis and others that restricted oxygen uptake, in association with high temperature, is the cause of normal dormancy in seeds and, indeed, in buds also. Thus, Vegis has pointed out that the embryos of developing seeds are liable to experience oxygen deficiency because of the surrounding seed coats and maternal tissues, and postulates that under such conditions of partial anaerobiosis normal oxidative breakdown through the tricarboxylic acid cycle and "terminal oxidation", necessary for growth in most species,

does not take place. Instead, the products of glycolysis, such as phosphoglyceric acid, cannot undergo normal oxidative breakdown, but become diverted into alternate pathways, leading to the formation of fatty acids and fats, which tend to accumulate in dormant tissues.

The possible importance of oxygen deficiency within the seed as a factor in dormancy is suggested by work on rice seeds, which are dormant immediately after harvesting, but gradually emerge from dormancy during dry storage. The dormant seeds of rice can be induced to germinate by removing the husks, thus indicating the importance of coat effects in these seeds. Storage in oxygen greatly reduces the dormancy period, suggesting that some oxidation reaction may be involved in the after-ripening processes during dry storage. However, Roberts tested the effects of various respiratory inhibitors (including inhibitors of terminal oxidation, Krebs cycle and glycolysis) and obtained the unexpected result that they *stimulated* germination of dormant rice seeds. He suggested that it is necessary for some oxidation reaction to proceed before germination can take place, and that this reaction is in competition with respiratory processes involving glycolysis, the Krebs cycle and the terminal oxidase system for the low levels of oxygen present in the seeds; hence the inhibition of these respiratory processes by various substances will release greater amounts of oxygen for the other oxidation reaction, which he suggests may involve the "pentose phosphate pathway" of carbohydrate metabolism. There is indeed evidence that the loss of dormancy is accompanied by a shift from the glycolytic to the pentose phosphate pathway in several species of seed. It has been further suggested that the action of respiratory inhibitors in stimulating germination is brought about by their inhibiting the enzyme catalase, which catalyses the breakdown of hydrogen peroxide. The hydrogen peroxide so spared is postulated to enhance the activity of the pentose phosphate pathway.

There is also some evidence that the effect of seed coats may be due to mechanical resistance to the growth of the radicle. Thus, several types of dormant seed will germinate if the seed coat is removed in the radicle region, but if seeds so treated are placed in a high osmotic concentration of 0.3 M mannitol (which will reduce the ability of the seeds to take up water and hence replace the mechanical effect of the seed coat) their germination is inhibited. However, the osmotic effect of the mannitol solution can be overcome by treatments, such as exposure to light or treatment with gibberellic acid, which will stimulate germination of the intact seed. It is concluded, therefore, that the normal effect of the seed coat is a mechanical one, to overcome which the radicle needs to develop a sufficient turgor. Whether this mechanical effect of seed coats is important in many species is not yet clear. Nevertheless, whatever the effect of the coats, it is clear that they play a very important role in many forms of seed dormancy.

SIMILARITIES BETWEEN SEED DORMANCY AND BUD DORMANCY

Inhibition of germination by the seed coats cannot be important in seeds which show embryo dormancy, where even the naked embryos are dormant. Hence we must seek

some other cause of dormancy in such cases. Now the dormancy of seeds showing a light or chilling requirement shows certain features in common with that of buds and other organs, which may be summarized briefly as follows:

- (1) Chilling for several weeks at 0-5°C is effective in breaking the dormancy of buds, rhizomes, corms and many types of seed.
- (2) Certain substances will break the dormancy of several kinds of organ; thus, thiourea and gibberellic acid will remove the dormancy of tree buds, potato tubers and several types of seed.
- (3) Certain tree buds and certain seeds may be induced to grow by exposure to long days, whereas short days are ineffective.

The close parallel between dormancy in buds and in seeds is particularly clear in instances where the buds and seeds of a single species are compared. For example, in birch (*Betula pubescens*) the dormancy of both the seeds and buds can be removed by chilling, by exposure to long days or by gibberellic acid. This parallel between seed and bud dormancy in a single species strongly suggests that the cause of dormancy is the same in both organs.

Now, bud dormancy is apparently not due to interference with gaseous exchange by the bud scales since (1) many dormant buds are not tightly enclosed by the scales, and (2) removal of the bud scales does not usually cause resumption of apical activity. Moreover, interference with gaseous exchange cannot be important in the *induction* of dormancy in buds, since until the buds are actually formed there can be no question of interference with oxygen uptake by the bud scales. On the other hand, we have seen that in many woody species resting buds are formed under short days, and that the response is determined by the daylength conditions to which the *leaves* are exposed. In view of the evidence for the role of hormones in the control of bud dormancy it is pertinent to examine their possible importance in some forms of seed dormancy.

HORMONAL CONTROL OF DORMANCY

Since hormones appear to play an essential role in most aspects of growth and differentiation it is reasonable to examine their possible role in the control of dormancy of both buds and seeds. Studies on this problem involve two main types of approach, viz. (1) observations on the effects of application of exogenous hormones, and (2) investigations on endogenous hormones, especially to establish whether there is any meaningful correlation between variations in the levels of endogenous hormones and the state of dormancy of buds and seeds.

Experiments with exogenous hormones have shown that the dormancy of many seeds can be overcome by application of gibberellins, cytokinins and ethylene. Species which show a response to gibberellins include a number which normally require after-ripening in

dry storage, others which are light-requiring and many which have a chilling requirement. Among the various gibberellins which have been tested, GA₄ and GA₇ have been found to be particularly active in stimulating germination of dormant seeds. In a smaller proportion of species dormancy can be overcome by cytokinins; usually a given species responds either to gibberellins or to cytokinins, but some seeds respond to both types of growth substance, e.g. lettuce, pear (*Pyrus communis*) and sugar maple (*Acer saccharum*). Ethylene has also long been known to stimulate germination in a number of species and in some species, such as lettuce, ethylene increases the germination percentage which can be obtained with cytokinins or gibberellins alone.

Although greatly increased dark germination can be obtained in many light-requiring seeds by application of exogenous growth substances, these do not always fully replace the light requirement. Thus, red light and gibberellins have often been found to be synergistic in overcoming dormancy, suggesting that their modes of action are not identical. Similarly, in certain weed species, such as *Spergula arvensis*, the effects of red light, ethylene and carbon dioxide are markedly synergistic in promoting germination. Again application of kinetin reduces, but does not entirely replace, the light requirement of lettuce seeds.

The dormancy of resting buds of many types of woody plants and other resting organs may be overcome by the same three main types of hormone as are active with dormant seeds. In most such cases the exogenous hormone removes a chilling requirement, but in buds of beech and birch exogenous gibberellin will also substitute a requirement for long photoperiods. It has been claimed that GA₃ is only effective in hastening bud-break in *Acer pseudoplatanus* after the chilling requirement has been met, but there seem to be other well-authenticated instances in which GA₃ is effective in replacing a chilling requirement, e.g., in unchilled hazel seeds.

Studies on changes in the levels of endogenous growth-promoting hormones in plant extracts have provided a number of instances in which levels of gibberellins and cytokinins have been found to decline during the development of dormancy and to increase during emergence from dormancy. Thus, it is well established that whereas levels of endogenous gibberellins and cytokinins are very high in young developing embryos, the levels of these hormones decline drastically to almost zero during the later stages of seed development (Fig. 11.7), although these changes are not confined to species which have dormant seeds. Concomitant with the decline in free gibberellins, there is an increase in gibberellin conjugates, especially gibberellin glucosides and glycosyl esters, which may serve as "reserves" in the seed. A similar decline in endogenous gibberellin and cytokinin levels appears to occur in the resting buds of woody plants during the development of dormancy, both under natural conditions and in response to short days applied experimentally.

By contrast, it has been found that levels of extractable gibberellins and cytokinins increase in both seeds and buds during chilling treatments. In several species of seed, the levels of cytokinins and gibberellins rise to a peak in succession and then decline, so that by the end of the chilling period these hormones have returned to a low level (Fig. 11.8). In several light-requiring seeds rapid increases in gibberellins and cytokinins have been observed following exposure to short periods of red light (Fig. 11.9). Thus, both chilling

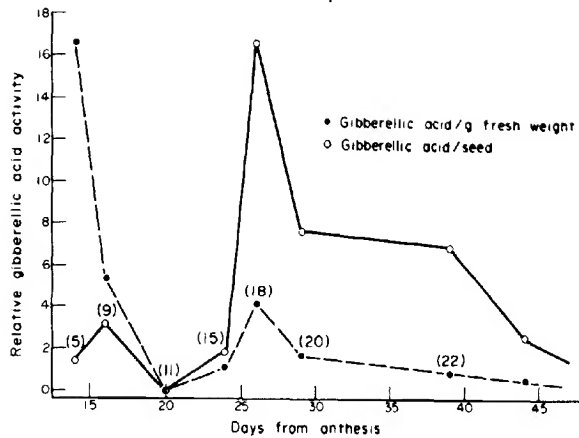


FIG. 11.7. Changes in endogenous gibberellin activity in seeds of *Phaseolus vulgaris*, during their development. Relative gibberellin activity per gramme fresh weight and per seed plotted against days from anthesis. The ordinate scale refers to gibberellin acid per seed; one unit of this scale is equivalent to 5 units of gibberellin activity per gramme fresh weight. (From K. G. M. Skene and D. J. Carr, *Austr. J. Biol. Sci.* **14**, 13-25, 1961.)

treatments and red light, which overcome the dormancy of various species of seed, result in increases in endogenous gibberellins and/or cytokinins.

We have seen that ethylene will stimulate germination in seed of a number of species, and there is increasing evidence that endogenous ethylene may play an important role in seed dormancy. Thus, the embryonic axes of non-dormant varieties of peanut (*Arachis hypogaea*) produce ethylene during germination whereas those of dormant varieties produce only low levels. In dormant *Xanthium* seeds there is a complex interaction between the effects of ethylene, carbon dioxide and oxygen in the promotion of germination. Non-dormant seeds actively produce endogenous ethylene under aerobic conditions, whereas dormant ones show only a low rate of ethylene production, so that dormancy in *Xanthium* seeds may be caused by the repression of the capacity for ethylene biogenesis. Germination of the dormant seeds can be stimulated by thiourea and the cytokinin, benzyladenine, which also increase ethylene production, but it would appear that the primary effect of these substances is to stimulate growth and that the increased production of ethylene is a consequence of such growth rather than its cause.

Although such general correlations between hormone levels and the state of dormancy suggest that the effects of dormancy-breaking treatments may be mediated through the variations in endogenous hormone levels, it still remains to be demonstrated unequivocally that this is the case, since (1) it remains to be shown whether the variations in the hormones are the cause or the effect of the changes in the state of dormancy, and (2) more

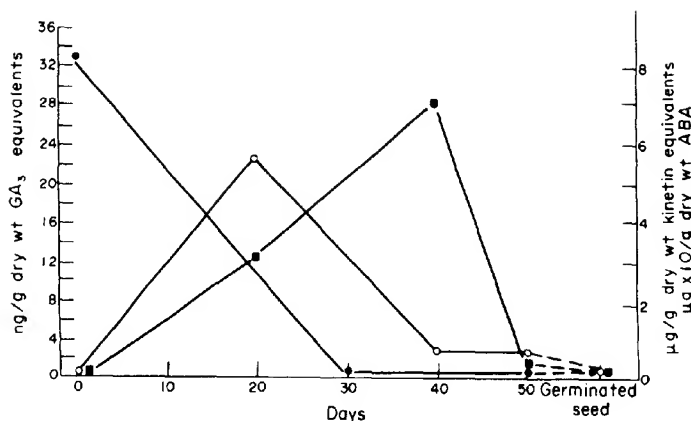


FIG. 11.8. Effects of chilling at 5°C on the levels of endogenous cytokinins, gibberellin-like substances and abscisic acid in seeds of *Acor saccharum*. Squares, acidic gibberellin-like substances. Circles, cytokinin-like substances. Solid circles, endogenous abscisic acid. (From D. P. Webb, J. van Staden and P. F. Wareing, *J. Exp. Bot.* 24, 105-16, 1973.)

detailed studies have sometimes shown a lack of a close correlation between levels of extractable hormones and dormancy. For example, although cytokinins increase rapidly in seeds of dock (*Rumex obtusifolius*) following exposure to red light (which stimulates their germination), there is evidence that these increases in endogenous cytokinins is *not* the primary cause of the release from dormancy. Moreover, most of the determinations of endogenous hormone levels in the past have been carried out on relatively crude plant extracts, using biological assay techniques, with all the errors inherent in these methods.

Apart from the problem as to the role of gibberellins, cytokinins and ethylene in the control of dormancy, there is good reason to believe that dormancy is not simply brought about by the absence of these growth-promoting hormones. On the contrary, there is circumstantial evidence to suggest that metabolism is actively blocked in dormant tissues, suggesting that possibly dormancy also involves natural growth-inhibiting substances. Substances which inhibit growth in various tests can be extracted from many plant tissues and hence it is possible that dormancy involves the active inhibition of growth by such substances, as was first suggested by Hemberg. He showed that extracts of dormant potato tubers and buds of ash (*Fraxinus excelsior*) contain substances which inhibit growth of *Avena* coleoptiles and that treatments, such as exposure to ethylene chlorhydrin, which overcome the dormancy of potato tubers also cause a marked reduction in the inhibitory activity of tissue extracts. It was also shown for a number of woody species that extracts of resting buds become less inhibitory during the course of the winter and this change is correlated with a gradual emergence of the buds from dormancy. However, again it is clear that such

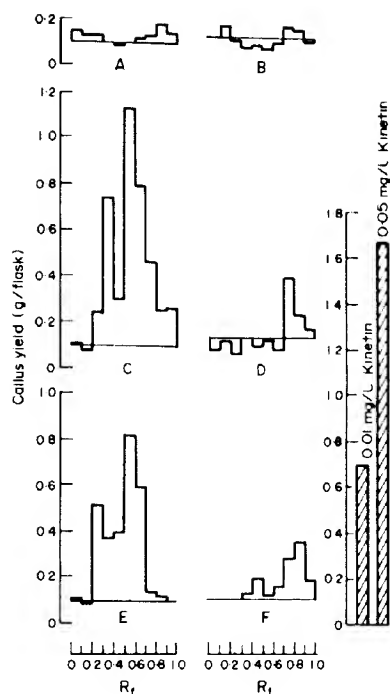


FIG. 11.9. Effects of short exposures to red (R) and far-red (FR) light on endogenous cytokinin levels in seeds of *Rumex obtusifolius*. All seeds were first imbibed in the dark for 2 days and then treated as follows before extraction of cytokinins: A, nil treatment (extracted immediately); B, 2 days in dark; C, 10 minutes R light; D, 10 minutes R light followed by 2 days in dark; E, 10 minutes in R light and 5 minutes FR light; F, 10 minutes R light and 20 minutes FR light. After paper chromatography seed extracts were assayed for cytokinin activity by the soybean callus assay. (From J. van Staden and P. F. Wareing, *Planta*, **104**, 126-33, 1972.)

a correlation does not establish a causal relationship between the level of growth inhibitors and dormancy. Bioassays can rarely differentiate between changes in amounts of "inhibiting" compounds and of growth-promoting compounds which interact with them. Moreover, because certain substances extracted from plant tissues inhibit growth of coleoptiles it does not necessarily follow that such substances normally function as growth inhibitors in the tissues from which they were extracted. It is probable, indeed, that some of these "inhibitors" are toxic substances which are normally restricted to the vacuoles of differentiated cells and hence do not normally have access to the growing tissues of the plant.

However, considerable interest was aroused by the discovery of abscisic acid (ABA), which is a powerful growth inhibitor in many tests. It was, indeed, the search for a natural dormancy-inducing substance ("dormin") which led to the isolation of ABA from leaves of *Acer pseudoplatanus* contemporaneously with its isolation from cotton fruits.

The application of exogenous ABA inhibits growth and germination in many tests. However, to inhibit the germination of non-dormant seeds, ABA must be supplied continuously, and if the seeds are rinsed in water they germinate rapidly. On the other hand, ABA induces the formation of resting buds ("turions") in the duck weed, *Spirodela polyrrhiza*, and the experimentally induced resting buds appear to show normal dormancy which can only be overcome by chilling or by application of cytokinins. Similarly, dormant immature embryos of yew (*Taxus baccata*) can be made to germinate by soaking them in a nutrient medium, which results in leaching of the endogenous ABA from the embryos. However, these leached embryos can be rendered dormant again by treating them with exogenous ABA. Again, it is possible to induce the formation of resting buds in seedlings of sycamore (*Acer pseudoplatanus*) by application of exogenous ABA. However, it is necessary to apply relatively high concentrations of ABA for prolonged periods in order to induce dormancy, so that it is not clear whether the formation of resting buds by this treatment is of any significance for the normal process. Moreover, some unsuccessful attempts to induce bud formation by application of ABA have been reported.

The effects of ABA and of gibberellins and cytokinins are mutually antagonistic in a number of tests. For example, the inhibitory effect of ABA on the germination of lettuce seeds can be completely reversed by cytokinins such as kinetin (Fig. 11.10). Similar interactions between ABA and GA_3 have been found in other tests. These observations have led to the idea that dormancy may be regulated by an interaction between growth inhibitors such as ABA, and growth promoters, such as GA_3 and cytokinins. In a number of seed species, the effects of ABA can be overcome by cytokinins, but not by GA_3 . It has been suggested that gibberellins, ABA and cytokinins have "primary", "preventive" and "permissive" roles, respectively, in the regulation of seed dormancy, i.e. it is postulated that gibberellins have a primary role in overcoming seed dormancy, that at appropriate concentrations ABA prevents germination, but that its effect can be overcome by conditions which lead to an increase in endogenous cytokinins.

Although experiments with exogenous ABA seem to point to its possible role in the regulation of dormancy, determinations of levels of endogenous ABA have given some results which appear to support the hypothesis, but others which appear to be inconsistent with it. Thus, endogenous ABA levels decline during chilling in seeds of apple and sugar maple (*Acer saccharum*), but experiments with other seeds have given less clear-cut results. Similarly, the ABA levels in buds of black currant (*Ribes nigrum*) and beech (*Fagus sylvatica*) have been reported to decline during the course of the winter, accompanied by a concomitant increase in the levels of the glucosyl-ester of ABA, suggesting a possible conversion of free ABA into conjugated forms. On the other hand, similar studies on birch and sycamore buds gave less marked changes in the levels of free ABA and the ratio of free/bound ABA, during the winter.

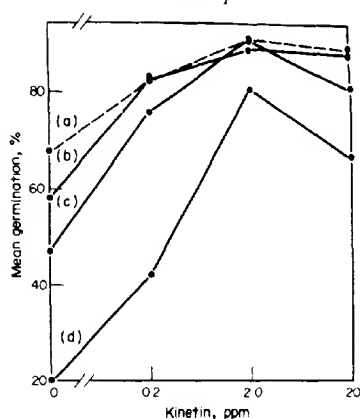


FIG. 11.10. Effect of abscisic acid and kinetin alone and in combination on the germination of lettuce seed. ABA concentration, ppm: (a) nil; (b) 0.002; (c) 2.0; (d) 20. (From P. F. Wareing, J. Good, H. Potter and J. A. Pearson, *S.C.I. Monograph* No. 31, Soc. for Chem. Ind., London, 1968.)

Since the formation of resting buds in woody plants is promoted by short days it might be expected, if bud formation and dormancy involves ABA, that levels of endogenous ABA would be higher under short days than under long days, but no such differences could be detected in the leaves and shoot apices of birch, maple and other species. Again, if high levels of ABA bring about cessation of growth and bud formation in woody plants, the very high ABA levels resulting from drought stress (p. 69) would be expected to be very effective in causing the formation of resting buds, but this does not appear to be the case in birch seedlings maintained in long days under water stress, although drought does promote bud formation in other species.

Thus, the present state of knowledge regarding the role of hormones in the control of dormancy remains confused, and although it seems very likely that variations in endogenous gibberellins, cytokinins and ABA are important in bud and seed dormancy, their precise role remains uncertain.

THE LONGEVITY OF SEEDS

The period during which seeds retain their viability varies greatly between species. The seeds of certain species remain viable for only short periods if kept in air at ordinary atmospheric conditions of humidity and temperature. Thus, seeds of willow (*Salix*) are viable for only a few days and must be sown very shortly after attaining maturity. The seeds of

poplar and elm retain viability for a few weeks, and those of oak, beech and hazel for a few months, when stored under cool, moist conditions.

The seeds of most species, however, remain viable for considerable periods, generally for at least a year and frequently for much longer. Various types of experiments have been carried out to determine the longevity of such seeds. Thus, various workers have tested the viability of very old seeds from herbaria, and Becquerel found appreciable germination of seeds of various members of the Leguminosae ranging from 100 to 200 years old. Other families characterized by long-lived seeds are the Euphorbiaceae, Malvaceae, Convolvulaceae, Solanaceae, Labiatae and Compositae.

Several long-term experiments have been set up in the United States to determine the longevity of seeds. In 1902, the U.S. Department of Agriculture set up an experiment in which seed of 107 species were placed in sterile soil in pots, which were then buried outside at different depths. After 20 years, some seeds of fifty-one of the species were still viable, but the seeds of most cultivated plants tested were dead. After 39 years, low germination was shown by twenty species and high germination by a further sixteen species.

A number of field observations also support the finding that seeds may retain their viability for long periods in the soil, under natural conditions. Thus, there is a well-authenticated case in which viable seeds of arable weeds were found in the soils of forests which had been planted on farm land some 20–46 years previously and had presumably lain dormant in the soil for that time. Other similar examples are known. Thus, living seeds of the Indian water lily, *Nelumbium incifera*, were found in the bed of a former lake in Manchuria, which was estimated to have been drained at least 120 years, and more probably 200 years, previously. Seed from herbarium specimens of this species which were 237 years old have also germinated. The seeds of *Nelumbium* have very thick coats and are impervious to water, so that the embryo is not imbibed with water until the coat is rendered permeable. Similarly, the hard-coated seeds of the Leguminosae will survive in the soil without taking up water until the coats have been eroded by the activities of soil micro-organisms. Such seeds may therefore lie for long periods in the soil in the *unimbibed* condition.

Many other types of seed which survive in the soil for long periods do not have impermeable coats, however, and hence they must survive in a moist condition. This fact raises the problem as to why it is that these seeds do not germinate in the soil, where they appear to have adequate conditions of moisture and temperature for germination. This problem has not been satisfactorily solved, but it has been suggested that high carbon dioxide concentrations in the soil render the seeds dormant. However, recent studies have shown that a very high proportion of buried weeds have a light-requirement for germination. It is of interest that among the buried seeds found to be light-requiring are those of certain species which do not normally show a light-requirement, and hence it would appear that burial in some way leads to the development of a light requirement. Where such light-requiring seeds lie buried in the soil they will remain dormant until ploughing or some other disturbance bring them to the surface, when they rapidly germinate. Seeds showing this behaviour include those of *Digitalis purpurea*, *Juncus* spp., *Polygonum* spp., *Veronica persica*, *Spergula arvensis* and *Hieracium* spp.

The ultimate cause of the loss of viability of seeds is not understood, but it is known that seedlings from old seeds show various cytological abnormalities such as chromosome breakage, disturbance of the mitotic spindle, etc., and physiological abnormalities such as chlorophyll deficiency. It is likely, therefore, that the loss of viability is due to slow chemical changes such as the denaturation of proteins.

GERMINATION

The Conditions for Germination

Since the tissues of the ripe seed are in a highly dehydrated condition, it is not surprising that water supply is frequently a limiting factor controlling the germination of seeds. Most seeds take up a relatively large amount of water when planted in a moist medium, and this is initially an imbibitional process, in which various substances present in the seed, especially proteins and starch, are involved. The imbibitional forces involved are enormous and certain seeds can take up considerable quantities of water from relatively dry soil.

Temperature is a second factor which plays an important role in controlling germination. The minimum temperature at which germination can occur varies considerably from one species to another, and the seeds of some species, such as beech, will germinate at temperatures only a little above freezing, whereas the seeds of tropical and sub-tropical species have much higher temperature requirements. Whereas most species will germinate under constant temperature conditions, other species require a daily alternation in temperature. For example, the seed of the dock, *Rumex obtusifolius*, germinates best when subjected to daily temperature alternations of 15° and 30° C. Alternating temperatures are also required by seeds of evening primrose (*Oenothera biennis*), Yorkshire fog grass (*Holcus lanatus*) and celery (*Apium graveolens*). Presumably these requirements are usually met by the normal variation between day and night temperatures under natural conditions. Little is known of the physiological basis of this phenomenon.

Although it is well known that seeds need conditions of adequate aeration for germination, there is little precise information on the oxygen requirements of seeds. It is clear, however, that the oxygen requirements of the intact seed will depend not only upon the metabolic demands of the embryo, but also on the permeability of the enclosing testa or other seed coats. As we have seen, these seed-coat effects appear to play an important role in the dormancy of many species. Whereas the oxygen content of the air is far above that needed for normal growth of plants, it is not far above that required for the germination of many seeds, no doubt due to the physical barrier to oxygen uptake presented by the seed coats.

There are marked differences between species in the ability of their seeds to germinate under water, no doubt because the partial pressure of the oxygen dissolved in water is considerably less than in air. Other seeds will germinate well under water; these latter include both aquatic species, and certain land species, such as rice.

The Process of Germination

In non-dormant seeds, active metabolism evidently commences soon after they are placed under conditions favourable for germination. The question arises as to what are the first stages in the complex overall process known as germination. Usually, of course, we take the emergence of the radicle as the primary criterion of germination, i.e. for most practical purposes the initiation of growth is taken as the first detectable sign of germination. It appears that the initial elongation of the radicle involves cell extension rather than cell-division, but cell division starts very early in the growth of the radicle. It is probable that the commencement of radicle growth is preceded by a number of preliminary processes, but little is known regarding the nature of these processes.

In the dry condition of the resting seed, metabolism must obviously be at an extremely low level on account of lack of water, but full metabolic activity does not develop immediately water is imbibed, even in non-dormant seeds.

When a non-dormant seed is planted under conditions favourable for germination there is a rapid increase in the respiration rate, which can be detected 2-4 hours after soaking in the case of peas. After this initial rise the respiration rate in peas reaches a steady value which is maintained for several hours. At about the time that the testa is broken by the radicle there is a further rapid rise in respiration rate, suggesting that in the initial phases of germination gaseous exchange is limited by the testa. By contrast, in barley and wheat there is a fairly uniform rise of respiration rate during germination.

The changes in carbon dioxide output (Q_{CO_2}), of oxygen uptake (Q_O) and of the respiratory quotient (RQ) during germination provide an indication of the type of respiration occurring (i.e. whether aerobic or anaerobic) and of the nature of the respiratory substrate, i.e. whether carbohydrate, fat or protein. During the early stages of germination of peas respiration appears to be predominantly anaerobic, owing to the restriction of oxygen uptake by the testa, and ethanol may accumulate in the tissues. The enzymes of glycolysis have been shown to be present in pea seeds.

The enzymes of the tricarboxylic acid (TCA) cycle by which aerobic respiration occurs are located in the mitochondria; it appears that the mitochondria in dry seeds are not fully active and are incapable of carrying out oxidative phosphorylation, but their activity increases during the later stages of germination. Also, it has been found that the electron transport system involved in terminal oxidation is not active in dry seeds. On the other hand, the pentose phosphate pathway, which provides an alternative mechanism for the aerobic respiration of carbohydrates, is active in bean seeds.

Although many enzymes are present in dry seeds, other enzymes are absent or present in an inactive form, and their activity only appears as germination progresses; examples of these are provided by several of the amylases, lipases and proteases responsible for the breakdown of reserve materials during germination. It has clearly been demonstrated that certain enzymes are synthesized *de novo* during germination. An excellent example is provided by the enzyme α -amylase, which is not present in the dry barley grain, but which appears during germination (p. 90). It is apparently secreted by the aleurone layer and

brings about the breakdown of starch in the endosperm. Normally the presence of the embryo is necessary for the appearance of α -amylase, suggesting that the production of the enzyme depends upon the supply of some substance from the embryo. It has subsequently been shown that the synthesis of α -amylase can take place in barley grains from which the embryo has been removed, if they are supplied with gibberellic acid, suggesting that the embryo normally supplies a natural gibberellin which initiates α -amylase synthesis in the aleurone layer. In this way enzyme synthesis is regulated, so that it does not take place until the embryo commences growth. It is now known that the synthesis of other enzymes, including RNA-ase and proteolytic enzymes, may be stimulated in barley grains by gibberellic acid.

Studies have been carried out on the RNA changes occurring in seeds during germination. It is found that there is active RNA synthesis during germination, affecting all fractions of RNA, and, in particular, there is marked increase in the monoribosome and polyribosome fractions. During the first 30 minutes of imbibition of water by wheat embryos there is a rapid formation of functional polyribosomes, with a corresponding increase in protein synthesis. Preparations of ribosomes from dry wheat embryos show little capacity to incorporate ^{14}C -leucine into protein, whereas preparations from imbibed embryos do so. However, if polyuridylic acid is supplied (as a synthetic messenger RNA), the ribosomes of dry seeds are found to be as active as those from imbibed seed. Thus, the inability of the ribosomes from dry seeds to carry out protein synthesis apparently does not lie in any deficiency in the ribosomes themselves, but is apparently due to the unavailability of m-RNA. However, there is evidence that m-RNA is present in dry seeds, but is apparently inactive or is spatially separated from the ribosomes. However, it appears that this pre-formed m-RNA becomes activated after the seed is allowed to imbibe water and is then able to support protein synthesis. Thus, the antibiotic, actinomycin D, which inhibits the production of m-RNA by the transcription of DNA in the nucleus, does not inhibit protein synthesis and polysome formation during early germination. In cotton seeds actinomycin D inhibits incorporation of ^{32}P into polysomes, but has no effect on the incorporation of ^{14}C amino acids into protein. On the other hand, inhibitors of protein synthesis (i.e. at the "translation" stage) do inhibit this process in germinating cotton seeds. These observations suggest that protein synthesis during the early stages of germination does not depend upon the production of new m-RNA at that time, but is directed by long-lived m-RNA already present and which was produced during the development of the seeds. Supporting evidence for this latter conclusion has come from studies on developing cotton embryos, in which it was shown that m-RNA for the synthesis of the enzyme protease is evidently present during the last 20 days of seed development, and yet synthesis of the enzyme does not occur, apparently because the translation process is inhibited, possibly by endogenous abscisic acid.

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Senescence and Abscission

IN COMMON with all multicellular organisms, higher plants are mortal and the life of the individual plant is ultimately terminated by death. Before the death of the whole plant has occurred, however, it is likely that there will have been earlier death of a number of its organs and tissues. As a result of the activity of the apical meristem, the upper part of the shoot shows a prolonged embryonic condition, while at the same time senescence and death is occurring in the older lateral organs, notably the leaves, flower parts and fruits. In many plants the death of organs has a pronounced seasonal character and there is a regular annual loss of part of the shoot system. In trees, this annual loss is mainly confined to the lateral organs mentioned, but in some herbaceous perennials, e.g. dock (*Rumex*), nettle (*Urtica*), bracken (*Pteridium aquilinum*), the whole above-ground part of the shoot may die each year. In annual species, the whole plant, except the seeds, dies after flowering and fruiting. Thus, the death of plant parts is a regular feature of the annual cycle of growth and is much more common in plants than in animals.

Death of an organ or of the whole plant is always preceded by the process of *senescence*, which may be regarded as the final phase in development that leads to cellular breakdown and death. We may conveniently distinguish between *organ senescence* and *whole plant senescence*. In most plants each leaf has only a limited life span so that as the shoot continues to grow in height, the older leaves at the base tend to senesce and die progressively (Fig. 12.1). This pattern of senescence has been described as *sequential senescence*, and it must be distinguished from the *simultaneous* or *synchronous senescence* of leaves of temperate deciduous trees, which is so conspicuous in the "fall". *Fruit senescence* is seen during ripening of both succulent and non-succulent fruits. The ripening of succulent fruits is a complex process which ultimately terminates in the senescence and decay of the tissues.

Before considering plant senescence it is necessary to distinguish between *monocarpic* species, which flower and fruit only once and then die, and *polycarpic* plants, which flower and fruit repeatedly. Monocarpic species include all annual and biennial plants and also a certain number of perennial plants which grow vegetatively for a number of years and then suddenly flower, fruit and die. Examples of this latter type of plant are seen in the "Century Plant" (*Agave*) and bamboo, both of which may grow vegetatively for many years, before

the single reproductive phase occurs. Thus, in monocarpic species, death of the whole plant is closely connected with reproduction and is evidently genetically determined to occur at this stage in the life cycle. By contrast, in polycarpic species, which include both herbaceous perennials and woody plants, death of the whole plant is not normally associated with reproduction and there is great variation among the different individuals of a given species with respect to the length of the life span. Thus, in dicotyledonous woody plants, which are all polycarpic, the individual tree normally lives for many years, and there is no universal life span characteristic of the species. Indeed, apart from accident and disease, there would seem no reason why a tree should not live indefinitely, although there would no doubt be mechanical problems when the branches become too heavy to support themselves. In monocarpic plants, on the other hand, death is destined to occur at a given point in the life cycle, and we may speak of *programmed death* in such plants.

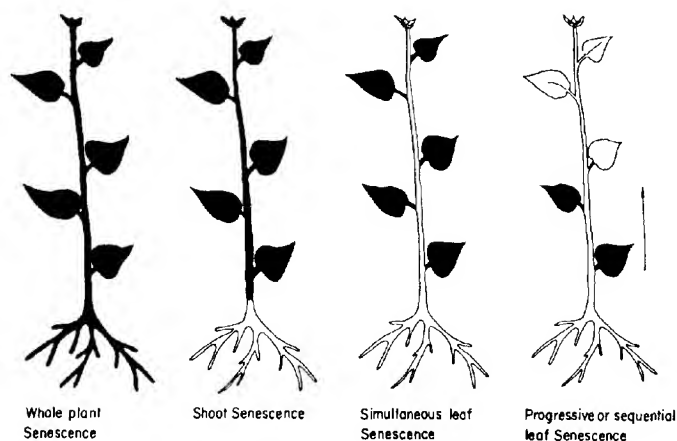


FIG. 12.1. Types of plant- and leaf-senescence.

What determines the life span of an individual cell, an organ or a whole plant? It might be suggested that a given cell has only a limited life span, which is determined by factors inherent in the cell itself. The loss of viability of seeds appears to be due to chromosome breakage within individual cells (p. 277), but it cannot be assumed that changes occurring in the dehydrated tissues of seeds in dry storage occur also in actively metabolizing cells and are a cause of senescence in the later stages of the plant cycle, although it has been suggested that chromosomal changes are important in the ageing and senescence of animal cells.

In some types of tissue, differentiation involves the early death of certain cells, such as those of vessel elements in the xylem, whereas neighbouring parenchymatous cells may

remain living for many years. The changes occurring in the protoplast during the differentiation of a vessel element may correspond closely to those occurring later in the constituent cells of a senescent organ, such as a leaf. However, the process of vacuolation and enlargement does not necessarily involve degenerative changes, since parenchymatous cells may live for many years, as in those of the pith and rays of some woody plants. Thus, it would seem that the maximum potential life of many types of differentiated plant cell is seldom reached in herbaceous plants, and that senescence and death do not occur on account of factors intrinsic in the individual cells, but because of conditions prevailing within the organ or organism as a whole. For example, sequential leaf senescence seems to be caused by competition between mature leaves and the growing regions of the shoot, and if a leaf is removed and allowed to form roots in the petiole it may live very much longer than if it had remained attached to the parent plant (p. 289). Thus the rate of senescence of plant organs is often under control of the whole plant and is not simply determined by intrinsic characteristics of the cells of that organ. However, it would appear that certain organs show inherent senescence processes, which are not under the control of the whole plant; thus, flowers and fruits undergo senescence whether they are allowed to remain on the parent plant or not.

In addition to the various "internal" factors of the plant which are involved in the regulation of senescence, a number of external factors may affect the rate of senescence, including drought, mineral nutrition, light intensity and daylength, and disease. We shall mainly be concerned with a consideration of the internal factors, but the importance of environmental factors must also be borne in mind.

THE BIOLOGICAL SIGNIFICANCE OF SENESCENCE

If we are correct in regarding senescence in monocarpic plants as "genetically programmed", then this implies that the process of senescence has arisen as a result of natural selection and that it has certain biological advantages to the species. What are these advantages? Why, for example, should it be any advantage to an annual species for the whole shoot to senesce during the development and ripening of the fruit --why should the leaves and other parts of the shoot not remain green, as they do during the ripening of the fruit in many polycarpic plants? The answer to this question probably lies in the fact that during the ripening of annual plants there is considerable breakdown of proteins to amino acids, which are exported from the leaves to the developing seeds and reutilized there as reserve material. For example, in the oat plant a large proportion of the nitrogen in the senescing leaves is transported to the fruits and accumulated there. Thus, recovery of nutrients from senescing organs constitutes a valuable saving to the rest of the plant. Similar export of reserve material occurs during the simultaneous senescence of the leaves of trees in the fall, the exported material being stored in the stem in this case. However, another advantage of leaf fall in deciduous trees probably lies in the resulting reduced rate of transpiration, which is probably essential for survival in climates in which the soil is frozen during the winter

(p. 254), while at the same time the return of leaf material and its breakdown in the surface litter releases mineral nutrients to the soil which are available for reutilization.

Sequential senescence of the basal leaves of a shoot not only releases reserves of nitrogenous and other substances for the young growing leaves, but may also bring about a saving of carbohydrates and other photosynthates where the basal leaves are heavily shaded and hence might actually become "parasitic" upon the plant by importing photosynthates from other parts which they would only consume in useless respiration. Thus, it is not difficult to see how senescence, as an active "programmed" process, may have several advantages.

THE MECHANISM OF SENESCENCE

Our present knowledge of the physiology and biochemistry of senescence is derived mainly from studies on leaves (both attached leaves undergoing sequential senescence and detached leaves or leaf discs) and on ephemeral flowers such as those of the Japanese Morning Glory, *Ipomoea tricolor*, the corolla of which opens and undergoes senescence within the space of only 24 hours (Fig. 12.10).

Sequential Leaf Senescence

The first visible sign of senescence is yellowing of the leaf, due to the breakdown of chlorophyll which renders visible the other leaf pigments, particularly the xanthophylls and carotenoids. A study of the fine structure of senescing leaves shows that there is progressive degeneration of the membrane structure of the grana of the chloroplasts, accompanied by the appearance of dense globules of lipid material (probably formed from the broken-down membranes) in which the carotenoid pigments dissolve. Other early changes involve the degeneration of the endoplasmic reticulum and the gradual disappearance of the ribosomes. The mitochondria retain their structure during the early stages of senescence, but later they undergo degeneration. The cells of the fully senescent bean leaf still retain an intact plasmalemma but the tonoplast disappears, and the structure of the cytoplasm and nucleus is almost completely lost, leaving the chloroplasts represented by vesicles containing lipid globules.

These structural changes in the cells of the senescing leaf are accompanied by changes in composition and metabolic activity. The protein content of the leaf declines progressively, as a result of the breakdown of proteins to amino acids and amides (Fig. 12.2). There is also a progressive decline in the RNA content of the leaf, with a particularly marked fall in ribosomal RNA (Fig. 12.3).

These degenerative changes are reflected in the rate of photosynthesis and respiration of the leaf. In *Perilla*, the rate of photosynthesis declines gradually from the time of full leaf expansion, accelerating during the later stages of senescence (Fig. 12.4). In this species the

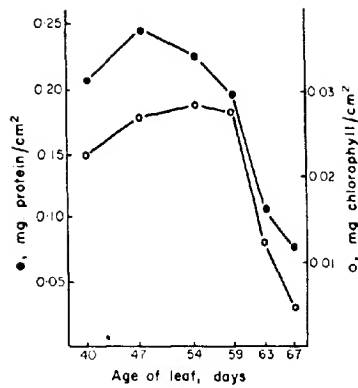


FIG. 12.2. Changes in protein and chlorophyll content of attached leaves of *Perilla frutescens* from expansion to abscission. (From H. W. Woolhouse, *Symp. Soc. Exp. Biol.* **21**, 179, 1967.)

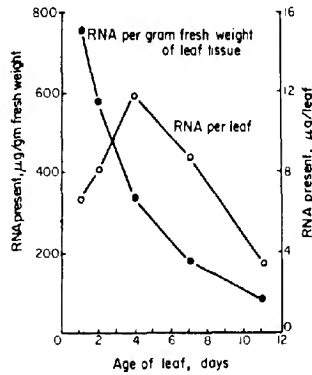


FIG. 12.3. Changes in ribonucleic acid (RNA) content of attached leaves of pea (*Pisum sativum*) from completion of leaf expansion. (From R. M. Smillie and G. Krotov, *Can. J. Bot.* **39**, 891, 1961.)

trend in photosynthetic rate follows closely the level of the soluble protein fraction of the chloroplasts known as "Fraction 1" containing the enzyme, ribulose-1,5-diphosphate carboxylase, which catalyses the process of carbon dioxide fixation. The trend in respiration rate seems to vary according to the species, but in some the rate appears to remain fairly constant until the final stages of senescence, when there is a sharp rise in respiration rate, corresponding to the "climacteric" observed during fruit ripening and senescence.

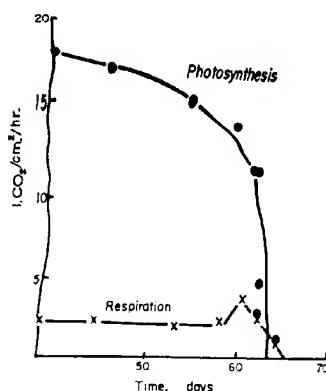


FIG. 12.4. Changes in rates of photosynthesis and respiration of attached leaves of *Perilla frutescens* from completion of expansion to abscission. (From H. W. Woolhouse, as for Fig. 12.2.)

The question arises as to what initiates and controls the degradative changes occurring during leaf senescence. The observation that, in some species at least, the respiration rate remains constant during the early stages of senescence suggests that it is not changes in respiratory metabolism which cause senescence. On the other hand, we have seen that a constant concomitant of senescence is marked decline in the protein and RNA contents of leaves, and close attention has been paid to these changes as possibly indicating the "key" processes in senescence. Now, it has been shown that a certain proportion of the leaf protein undergoes continuous "turnover", i.e. the protein is being continuously synthesized and broken down, so that the overall rate of change in protein content represents the net differences in the rates of these two processes. Where there is such a continuous turnover, a decline in protein content may reflect a fall in the rate of synthesis or a rise in the rate of breakdown, or both.

The breakdown of protein is brought about by proteolytic enzymes. Studies on changes in proteolytic enzyme activity have given no indication of increased activity during leaf senescence, and hence it would appear that the decline in protein content is primarily due to a reduced rate of synthesis. One possibility that has been suggested is that the senescent leaf retains its full capacity for protein synthesis and that the rate of synthesis is limited by lack of amino acids in the leaf. In a healthy, green leaf the amino acids released by protein breakdown are reutilized in further protein synthesis. But it has been suggested that in a senescent leaf amino acids are exported to other parts of the plant so rapidly that there is no "pool" of free amino acids available for protein synthesis, so that there is a decline in protein content (Fig. 12.5). It is, in fact, found that there is no appreciable accumulation of free amino acids in senescent attached leaves, no doubt due to continuous export. However, it is then necessary to explain why the supply of amino acids should be limiting in a senescent leaf and not in a normal one.

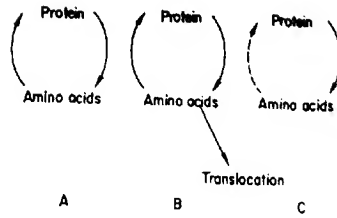


FIG. 12.5. Diagram to illustrate (A) turnover of leaf protein, (B) the translocation hypothesis, and (C) the hypothesis of a defect in protein synthesis. (From E. W. Simon, *Symp. Soc. Exp. Biol.* **21**, 215, 1967.)

Now, there is evidence that the sequential senescence of leaves arises from competition for metabolites and nutrients between old leaves at the base of the stem and young growing leaves in the apical regions. This conclusion is indicated by the fact that if a plant is decapitated and the axillary shoots are removed then the senescence of the older remaining leaves is greatly retarded and indeed leaves which are already showing signs of senescence may undergo a recovery and become green again, even if they were previously showing yellowing. The uppermost leaf of a decapitated plant may also undergo considerable growth and become abnormally large. Thus, it would appear that sequential senescence is a "correlative phenomenon", and shows resemblances to apical dominance (p. 129). Sequential senescence is more pronounced under conditions of mineral nutrient deficiency—for example, plants grown in too small pots frequently show marked senescence of the basal leaves, under which conditions there is presumably severe competition between young and old leaves for available nutrients, and in this competition the young leaves are evidently at an advantage. However, the senescence of the older leaves can be retarded or reversed by the application of nitrogenous nutrients, such as ammonium nitrate.

Thus, it is possible that the competition between young and old leaves results in a rapid rate of transport of amino acids from old to growing leaves and hence to a lower pool of these metabolites being available for protein synthesis within the old leaves. It is an essential part of this hypothesis that there should be active protein turnover, but in fully expanded *Perilla* leaves the "Fraction 1" protein is found to have zero turnover rate, i.e. there is no continuous synthesis and breakdown, and yet during senescence Fraction 1 declines more rapidly than a second fraction "Fraction 2", which does show active turnover.

Another postulate of the hypothesis we are considering is that the capacity for protein synthesis remains relatively unimpaired during leaf senescence. A convenient method for measuring the rate of protein synthesis is to determine the rate of incorporation of radioactive amino acids, such as ^{14}C -leucine, into protein. Similarly, the rate of RNA synthesis can be followed by the rate of incorporation of an RNA precursor, such as ^{14}C -adenine. Studies of this type have shown that the capacity of tobacco leaves to incorporate ^{14}C -leucine and ^{14}C -adenine declines during senescence, although quite yellow leaves retain some capacity to synthesize certain enzymes, such as peroxidase and ribonuclease (which

brings about the breakdown of RNA). It might be argued, however, that the decline in capacity for protein synthesis is the *result*, rather than the cause of senescence. Overall, nevertheless, it seems that protein metabolism in senescing attached leaves may be viewed as an unbalanced turnover reaction, with catabolism exceeding anabolism.

Thus we see that some evidence supports the hypothesis that the decline in protein content of attached senescent leaves is due to limiting levels of amino acids, rather than to a reduced capacity for protein synthesis, but that the evidence is not conclusive. Furthermore, it is known that when the young leaves are removed from a plant, the remaining older leaves "regreen" and show a marked and rapid rise in RNA synthesis which precedes a rise in leaf protein level. Within 12 hours of removal of the young leaves from *Perilla* plants the incorporation of $^{32}\text{PO}_4$ into chloroplastic ribosomal RNA is stimulated in older leaves, and this change occurs before an effect is seen in the cytoplasmic ribosomes of the leaf. Results such as these suggest that very early events in leaf senescence involve changes in nucleic acid metabolism within the chloroplasts.

We shall now consider an alternative approach to the problem of leaf senescence, in which use has been made of leaves, or portions of leaves, which have been isolated from the parent plant. Excision of a leaf usually results in the immediate onset of senescence processes in its tissues, and detached leaves or leaf discs therefore provide convenient material for experiments under controlled conditions, without the additional complication of correlative influences from other parts of the plant.

Senescence of Detached Leaves

So far, we have considered the natural senescence of leaves still attached to the plant, but it has been known for many years that when a green leaf is detached from its parent plant it rapidly deteriorates and shows signs of accelerated senescence.

As with attached leaves, the visible signs of senescence are accompanied by a decrease in the protein and RNA contents of the leaf. Protein breakdown commences remarkably soon after the leaf has been detached; for example, protein breakdown is detectable 6 hours after excision of barley leaves. The breakdown of protein commences at the same rate whether the leaves are kept in the light or the dark, but the rate of breakdown later decreases in the light, whereas it continues at a high level in the dark (Fig. 12.6). The initial equal rates of breakdown in light and dark suggest that senescence is not triggered off by carbohydrate deficiency, although this factor may be important during the later stages of senescence in the dark.

In detached leaves, protein degradation leads to the accumulation of amino acids and amides in the leaf, since they cannot be exported, as is the case with attached leaves, although there may be accumulation at the base of the petiole where this is present. Thus, senescence can occur in detached leaves, or in leaf discs, even though there is a high level of amino acids within the tissues. Hence, under these conditions the reduced protein synthesis cannot be attributed to lack of amino acids. On the other hand, detached leaves show reduced capacity

for protein synthesis, as indicated by the reduced capacity to incorporate ^{14}C -leucine into protein. Indeed, leaf discs which have remained in the dark for several days lose this capacity completely. Thus, there seems no doubt that the decline in protein content observed in detached leaves arises from a reduced capacity for protein synthesis.

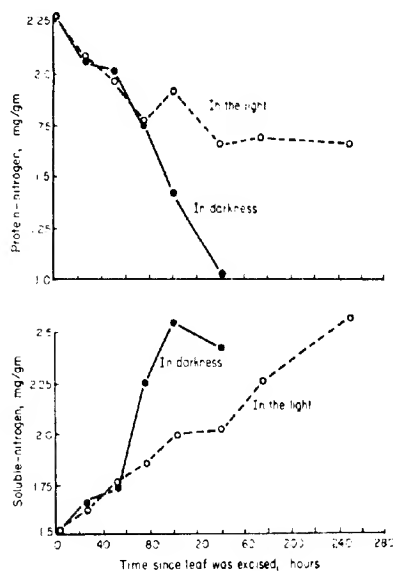


FIG. 12.6. Changes in protein and soluble nitrogen in detached tobacco leaves. Following excision, protein-N declines and soluble-N increases. Leaves kept in darkness show these changes to a greater extent and are yellow and senescent after approximately 100 hours. (From H. B. Vickery et al., *Connecticut Agric. Exp. Sta. (New Haven) Bull.* **374**, 557, 1935.)

It was observed that the rapid rate of protein breakdown in detached leaves is arrested if they are allowed to form roots (Fig. 12.7). Indeed, leaves which have been planted in soil and induced to form roots on the petiole will live for considerable periods. On these grounds, Chibnall suggested that roots must supply some "factor" which is necessary for the maintenance of protein synthesis in the leaves, but the nature of this "root factor" remained unknown. However, it was later found that kinetin will prevent the senescence of detached leaves if applied to them in aqueous solution. Thus, if a drop of kinetin solution is applied to a senescing tobacco leaf, then the area of leaf which received the kinetin will remain green, although the surrounding leaf tissue continues to yellow (Fig. 12.8). Similarly, if discs of radish or *Xanthium* leaves are placed on a solution of kinetin they still retain their green colour after the control discs (on water) have become fully senescent. Moreover,

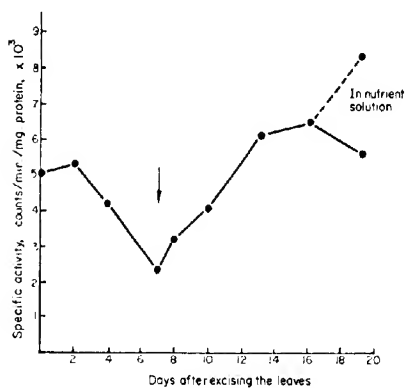


FIG. 12.7. Changes in capacity for protein synthesis (as measured by incorporation of ^{35}S -methionine into protein) in leaves of *Nicotiana rustica*. Note initial decline in capacity for protein synthesis, followed by recovery when roots appeared on petiole (indicated by arrow). (After von B. Parthier, *Flora, Jena*, **154**, 230, 1964.)

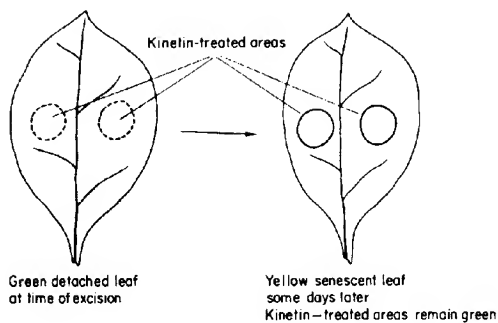


FIG. 12.8. Effects of kinetin on leaf senescence.

electron microscopic studies show that the kinetin-treated discs still have the normal structure of a green leaf.

Thus, senescence in detached leaves can be prevented either (1) by the presence of roots, or (2) by application of kinetin and other synthetic cytokinins. It was, therefore, natural to consider whether the natural "root factor" which delays senescence is an endogenous cytokinin. It is of great interest, therefore, to discover that cytokinins are present in the root exudate from sunflower, grape and other plants, and this suggests that leaves may depend

on the supply of endogenous cytokinins from the roots, for the maintenance of a normal, green condition. We shall return to this question later (p. 296).

Not only do cytokinins delay senescence of detached leaves, but they also cause a rapid increase in the rate of RNA and protein synthesis only a few hours after their application (Fig. 12.9); incorporation of precursors, such as ^{14}C -adenine or ^{14}C -cytidine, into all fractions of RNA appears to be increased by kinetin. Since protein synthesis involves the RNA "machinery" of the cell, the effect of cytokinins on protein synthesis is readily understood if their primary effect is upon RNA synthesis, but they may also delay senescence through their effect on the mobilization of metabolites (p. 295). The possible mode of action of cytokinins in stimulating RNA synthesis was discussed in an earlier chapter (p. 92).

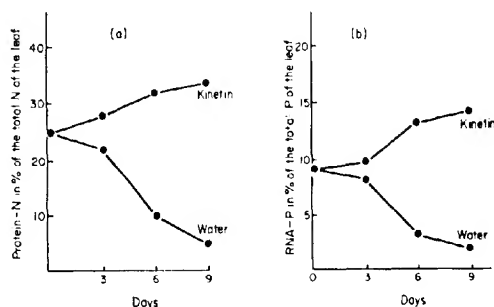


FIG. 12.9. Changes in (a) protein and (b) RNA content in kinetin- and water-treated halves of undivided leaves of *Nicotiana rustica*. (After R. Wollgast, *Flora, Jena*, **151**, 411, 1961.)

With most species, gibberellins do not affect the rate of senescence of leaf discs, but in *Taraxacum*, *Rumex*, *Tropaeum* and a few other herbaceous species, gibberellins are more effective in delaying senescence than cytokinins. On the other hand, auxins are found to be relatively ineffective in delaying senescence in the leaves of herbaceous plants, but they are active in this respect in the leaves of some deciduous woody perennials (e.g. cherry) and in pericarp tissues of dwarf beans (*Phaseolus vulgaris*). Applications of gibberellins may also delay the onset of senescence in leaves of some deciduous trees.

Contrasting with the senescence-delaying effects of cytokinins, gibberellins and auxins, both abscisic acid (ABA) and ethylene appear to accelerate the processes of senescence in leaves. Exposing leaves on intact plants, or detached leaves or leaf discs, to ethylene gas or to ABA usually results in accelerated yellowing. Fruit senescence (or ripening) is similarly speeded up by ethylene, and it is well established that ethylene plays an important role in the natural ripening of succulent fruits. Both ABA and ethylene inhibit RNA and protein synthesis in isolated leaf discs of many species, which emphasizes that their effects in senescence are opposite to those of cytokinins and gibberellins.

Senescence in Flowers

The evanescence of flowers is proverbial, and is normally a consequence of senescence in the corolla. In recent years, studies of the biochemical and physiological bases of flower senescence have tended to concentrate upon certain shortlived, or ephemeral, flowers such as those of the *Ipomoea tricolor* (Fig. 12.10). As also seen in leaves, it has been found that senescence in these flowers is accompanied by rapid falls in the levels of protein, RNA and DNA (Fig. 12.11). Exposure of newly opened flowers to ethylene causes petal fading and other senescent changes within 90 minutes, and it has been found that endogenous ethylene production rises at the same time that natural fading occurs and RNase levels rise.

The time-course of decreases in protein, RNA and DNA levels can be compared with changes in the levels of activity of hydrolytic enzymes such as proteinase, RNase and DNase (Fig. 12.11), and it can be seen that the breakdown of nucleic acids occurs at the same time that nuclease activity increases dramatically whereas the level of proteinase is practically unchanged throughout senescence. It seems clear that the decrease in protein levels which occur during senescence is not brought about by changes in total proteinase activity present in the cells, but rather by some process which allows proteinases to act on the cell proteins. The senescing cells of the corolla, like those of senescing leaves, retain the capacity for synthesis of certain proteins (e.g. the RNase that rises in activity during senescence is known to be synthesized *de novo*) and overall loss of protein therefore reflects an imbalance in protein turnover, as referred to previously (p.288), with catabolism exceeding anabolism.

There is accumulating evidence that many of the hydrolytic enzymes present in plant and animal cells are normally compartmentalized, in that they are located in particular regions and kept separate from other cellular constituents. This prevents the occurrence of *autolysis* (uncontrolled selfdigestion) in the cell. The regions, or compartments, which contain these hydrolyses are grouped together under the general term, the *lytic compartment*, which includes organelles such as *lysosomes*. As long as the lytic compartment remains intact, breakdown of cellular components is under control and is known as *autophagy*. The phenomenon of autophagy probably involves transport of substrate molecules across lytic membranes to come into contact with the hydrolases, followed by transport of hydrolysis products out of the lytic compartment. Because metabolism continues in senescent tissues, including protein synthesis, the lytic compartment is probably intact during most of the senescence process. The final step in senescence does, nevertheless, appear to be that of autolysis, involving rupture of the membranes of the lytic compartment.

Whole Plant Senescence

As we have seen, in monocarpic species, such as wheat (*Triticum*), soybeans (*Glycine max*) and French beans (*Phaseolus vulgaris*), senescence of the whole plant is usually associated with



FIG. 12.10. Development of the ephemeral corolla of *Ipomoea tricolor*. A. Mature flower bud in the early morning of the day of flowering. B. Anthesis at 6 a.m. C. Rigid ribs responsible for the shape of the funnel. D. First signs of fading and wilting in early afternoon. E. Partially rolled-up funnel at 5 p.m. F. Corolla in the morning of the day after flowering. (From Ph. Matile, *The Lytic Compartment of Plant Cells*, Springer-Verlag, Wien and New York, 1975. Original prints kindly provided by Professor Dr. Ph. Matile.)

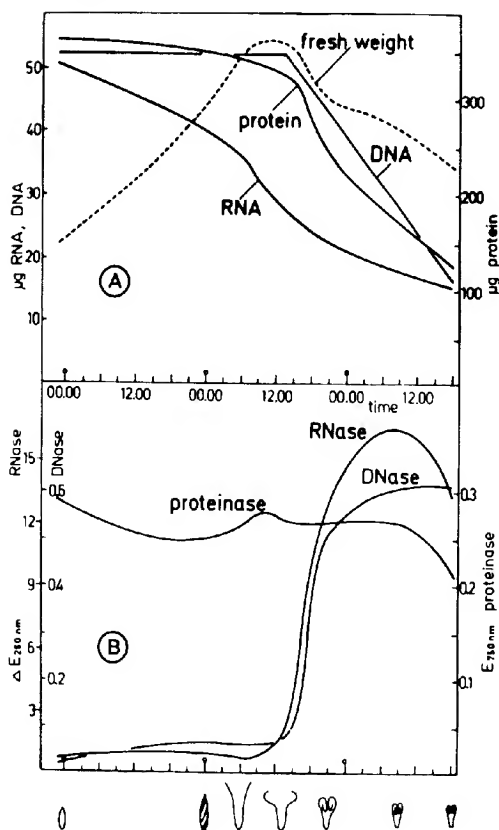


FIG. 12.11. Senescence in the ephemeral flowers of *Ipomoea tricolor*. Top: contents of protein and nucleic acids. Bottom: relative activities of proteinase and nucleases per corolla. The sketches below the time axis indicate the changing shape of the corolla—see also Fig. 12.10. (From Ph. Matile, *The Lytic Compartment of Plant Cells*, Springer-Verlag, Wien and New York, 1975. Original print kindly supplied by Professor Dr. Ph. Matile.)

the development of fruits. Thus, in the French bean plant, as the pods and seeds grow and approach their full size, there is perceptible yellowing of the leaves. When fully developed, the fruits yellow and ultimately lose water as they ripen, and at the same time the leaves and stems become progressively more senescent and in due course they die. Not only are plant senescence and fruit development closely correlated in time, but there does indeed appear to be a causal relationship between the two processes, as shown by the fact that removal of

all the flowers and young fruits from bean plants greatly delays senescence of the remainder of the plant. Moreover, the same effect is observed if, instead of removing the whole fruits, only the seeds are removed surgically from the pods. Thus, the presence of developing seeds appears to control plant senescence, suggesting that some "signal" is sent out from the developing seeds, which regulates senescence in other parts of the plant.

Now, the developing seeds of bean increase steadily in dry weight, and food reserves, including protein and starch, accumulate in the developing cotyledons. It is known that, at the same time, the breakdown of proteins and carbohydrates is occurring in the leaves and that amino acids, sugars and other metabolites are transported to the developing fruits. Similar effects are seen clearly in other species. It was, therefore, suggested by Molisch that developing seeds cause senescence in other parts of the plant by mobilizing and accumulating not only carbohydrates but also amino acids and other substances derived from the leaves.

The accumulation of reserve materials into fruits, tubers and other storage organs is well known. Similarly, actively growing meristematic regions, such as young leaves, are known to be able to mobilize nutrients from other parts of the plant. Such centres of mobilization are often referred to as "sinks" for nutrients. The manner in which mobilization by such sinks is achieved is not understood, since the mechanism of phloem transport is itself still not understood. However, in recent years increasing evidence has appeared indicating that growth hormones may play an important role in mobilization effects. Thus, when kinetin is applied to a small area on a senescing tobacco leaf, it is found that radioactive amino acids applied to the other parts of the leaf are accumulated at the point of application of kinetin. Thus, metabolites appear to be "attracted" towards regions which have been treated with kinetin. It is possible that this effect of kinetin on the mobilization of metabolites is a secondary one, arising from the stimulation of protein synthesis at the point of kinetin application, so that a metabolic "sink" is created, which in turn causes movement of metabolites towards this point. However, Möthes was able to show that if aminoisobutyric acid (which is not a naturally occurring substance and is not incorporated into proteins) is applied to a part of the leaf away from the point of kinetin application, it accumulates in the latter region, just as the natural amino acids do. Thus, on this evidence it would seem that kinetin may affect the movement of metabolites independently of any effect it has on protein synthesis.

There is evidence that auxin-directed transport may also play an important role in the movement of metabolites in ripening bean plants. We have seen (p. 132) that metabolites tend to accumulate in regions of high auxin content and it is known that developing bean seeds are rich sources of auxin. Thus, it is possible that the movement of reserve materials from the leaves into the developing seeds may be an example of auxin-directed transport. Evidence in support of this hypothesis has been obtained in the following experiment. The fruits were removed from young bean plants, and a decapitated peduncle (fruit stalk) was allowed to remain on each plant. In some of the plants indole-acetic acid (IAA) in lanolin was applied to the peduncle stump and in other plants (controls) plain lanolin was applied. It was shown that when radioactive phosphorus, ^{32}P (in the form of orthophosphate), was

applied to the stem bases, it very rapidly moved up into the peduncles to which IAA had been applied, but very little moved into the peduncles to which only plain lanolin was applied.

The hypothesis that plant senescence is brought about by the mobilization of nutrients by developing seeds would seem to be consistent with many observations, and if we postulate that the leaves become depleted of amino acids as a result of their export to the seeds, we have essentially the same hypothesis as was suggested above for the sequential senescence of leaves. However, there are certain observations which are difficult to reconcile with this hypothesis. Thus, it is found that in spinach (*Spinacia oleracea*), which is a dioecious species (with separate male and female plants), senescence of the male plants follows flowering, just as in female plants, although such male plants do not, of course, carry any developing fruits; moreover, removal of the male flowers delays senescence of the leaves. Further, in an experiment with *Xanthium pensylvanicum*, it was found that if all buds were removed from plants before exposure to short days, so that no flowers could be formed, the leaves of these plants later senesced at the same time as did those of plants which had been allowed to form flowers and fruit. On these and other grounds, the hypothesis that plant senescence can be explained simply in terms of the mobilization of nutrients by developing seeds is rejected by many workers.

Since protein synthesis is controlled by the RNA apparatus of the cell, it is entirely possible that the presence of developing seeds brings about protein breakdown in the leaves through an effect upon their RNA metabolism. For example, some fractions of RNA undergo rapid turnover, and it may be that certain RNA precursors are exported from the leaf to the developing seeds, and so are not available for reincorporation into RNA in the leaf. A different explanation has been suggested, as follows. We have seen that leaves undergo rapid protein and RNA breakdown when they are separated from the plant and it has been suggested that they are dependent on a continuous supply of cytokinin from the roots for normal RNA metabolism. Now, seeds are found to be rich in cytokinins. It is not known whether the total cytokinin content of seeds is synthesized there, or whether cytokinins are mobilized there from other parts of the plant. If the latter were the case, then it is possible that cytokinins produced in the roots are directed away from the leaves in the presence of developing seeds, with the result that the normal maintenance of the RNA apparatus of the leaf is not possible and hence protein synthesis is prevented, as in a detached leaf. One observation which is against this latter hypothesis is the fact that the senescence of attached leaves cannot normally be arrested by application of kinetin, suggesting that they are not deficient in cytokinin and hence their senescence must be due to some other cause. Thus, it would seem possible that the causes of senescence in a detached leaf are not the same as those of an attached leaf which undergoes sequential senescence or in response to fruiting.

It will be clear that although a number of interesting approaches to the problem of senescence in plants are being made, it is too early to be able to present a single overall hypothesis which will account for all the facts. However, in general one can recognize that the initiation of senescence involves an imbalance in the relative levels of growth hormones,

and that this change in hormonal status may be caused by either an environmental stimulus (such as short days) or by internal factors such as competition between spatially separated organs of the plant. Once set in motion, senescence involves autophagy and eventually autolysis.

SYNCHRONOUS LEAF SENESCENCE

The synchronous senescence of leaves seen in deciduous woody plants in the autumn or "fall" is so striking that it has given its name to this season of the year, in America at least. This type of leaf senescence differs rather markedly from sequential senescence in two respects. Firstly, it is primarily controlled by environmental rather than "internal" factors, such as competition between young and old leaves. Secondly, it appears to involve rather different hormonal factors.

Two environmental factors appear to be involved in determining the onset of senescence in deciduous woody plants, namely, daylength and temperature. It has long been known that short days tends to promote leaf senescence in woody plants such as *Liriodendron tulipifera* and *Ailanthus altissima*. Moreover, there have frequently been reports of delayed leaf fall in trees growing near street lights, so that they were exposed to long days as the natural daylength shortened. However, if seedlings of woody plants are rendered dormant by placing them under short-day conditions in a warm greenhouse, many species are found to retain their leaves in a green, healthy condition for several weeks at least. Thus, it seems likely that normal leaf fall is determined by short days in association with the low temperatures occurring in the fall, but precise experimental data on this matter are still lacking.

Although environmental factors are very important in controlling synchronous leaf senescence, influences from the rest of the plant are evidently also operating, since if a disc is nearly cut out of a cherry leaf by means of a cork borer, leaving only a small connection with the rest of the lamina, this disc remains green long after the remainder of the leaf has become senescent. Thus, it would seem that synchronous, as well as sequential senescence, depends upon the export of materials from the leaf.

We have seen that the senescence of excised leaves or leaf discs of herbaceous plants can be delayed by kinetin, and sometimes by gibberellin, but not by auxins. In woody plants, however, the reverse appears to be true. Thus, if a drop of the auxin, 2,4-D, is applied to cherry leaves in the fall, the areas receiving the auxin remain green long after the remainder of the leaf has become yellow. Similar results may be obtained with detached leaves of woody plants at all times of the year. Gibberellin will delay senescence of leaves of ash (*Fraxinus excelsior*). Just as application of kinetin and gibberellin enables the leaves of herbaceous plants to maintain their capacity for RNA and protein synthesis, so does 2,4-D for leaves of woody plants. Thus, it would seem that endogenous auxins probably play an important role in the natural senescence of leaves of trees. It is very likely that the influence of daylength on leaf fall is exerted through its effect on endogenous hormone levels, since there are lower levels of endogenous auxins and gibberellins in the leaves of woody plants under short days than under long days (Chapter 11).

We know little about factors controlling leaf senescence in evergreen broad-leaved trees and conifers. Some evergreen species may retain their leaves for several years. In some cases senescence of the older leaves occurs when a new suite of leaves is put out, suggesting that competition between old and new leaves may be important in such species.

HORMONES AND ABSCISSION

Shedding of both vegetative and reproductive organs and tissues is a naturally occurring process in all plants. Parts of plants are lost either by a process of death and withering, as in most monocotyledons and in plants such as potato with subterranean organs of perennation, or by the formation of abscission layers. The organ or tissue lying distal to the abscission layer falls to the ground by its own weight or by the action of an external force such as wind.

Abscission, or separation, processes are responsible for the shedding of tissues such as those of the bark of trees and the periderm in roots. However, because the loss of such tissues is usually continuous but slow, the phenomenon does not lend itself readily to experimental studies. In contrast, the abscission of leaves, flowers and fruits tends to be much more rapid and dramatic, and is often of importance in agriculture and horticulture. These facts have resulted in very much more attention being devoted to gaining an understanding of how abscission is achieved and regulated in such organs.

Leaf Abscission

In most plant species there comes a time during the life of each leaf when it is shed from the stem. This occurs most obviously at the end of the growing season in temperate regions of the world, but leaf fall is not confined to autumn. All through the summer in temperate areas, and all through the year in the tropics, there is a less conspicuous but continuous dropping of older leaves from plants. The process by which leaves (and also other organs such as fruits and flowers) are removed from the plant is known as *abscission*. The act of abscission of an organ, such as a leaf, is usually achieved by the formation of a *separation* or *abscission layer* at the base of the petiole. This is a thin plate of cells oriented at right angles to the axis of the petiole (Fig. 12.12). The walls of the separation layer cells become softened and gelatinous, so forming a weak region, which readily breaks under the strain of wind-induced movement of the leaf.

It is a general feature of abscission that, before the shedding process occurs, the organ that is to be discarded undergoes senescence. Senescence in a plant organ, such as a leaf, normally commences in its distal parts and spreads to the proximal regions. The last cells of the organ to undergo senescence are those situated immediately adjacent to the abscission zone, and at this stage the separation point is often clearly visible as a yellow/green junction. This junction therefore represents an interface between two physiologically dissimilar tissues,

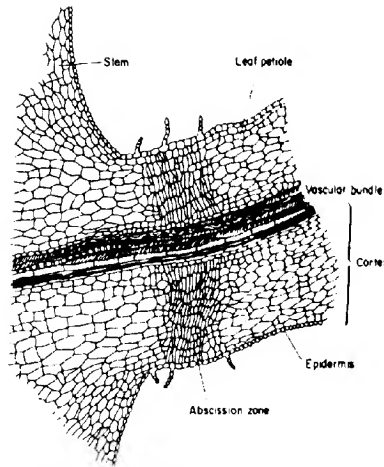


FIG. 12.12. Longitudinal section through the abscission region of the base of the petiole of a leaf of a typical dicotyledonous plant. (From J. Torrey, *Development in Flowering Plants*, Macmillan, New York, 1967.)

and it is the development of this interface at the abscission zone which signals the start of the abscission process.

To begin to understand the process of abscission it is necessary to identify the particular characteristic of senescent tissue which initiates the biochemical events of abscission. Over the past 10 years it has become increasingly clear from research that it is the relatively high levels of ethylene produced by the senescent cells distal to the abscission zone which is the important regulatory factor. The ethylene generated in the senescing tissues initiates the separation process in the abscission zone cells. In many plant species the whole of a senescent leaf produces large amounts of ethylene, whereas in others only senescing tissues of the petiole show a rise in ethylene production. In either case, however, the effect is to increase the ethylene concentration at the site of the abscission zone in the petiole.

Abscission of leaves can be artificially induced by exposing plants to ethylene at a concentration as low as $0.1 \mu\text{l l}^{-1}$, but the leaves of different ages vary considerably in their sensitivity to the gas. The oldest leaves are the first to abscise, with the remainder being shed sequentially so that the youngest are the last to fall. This sequential pattern of abscission response to exogenous ethylene appears to be related to variations in auxin levels in leaves, but the mechanisms by which auxin affects senescence and ethylene-sensitivity are not yet understood.

Many of the experiments conducted on leaf abscission have involved the use of isolated excised abscission zones (usually a nodal explant bearing petioles of each of which the lamina has been removed). With such an experimental system it has been found that application of an auxin, such as IAA, to the cut end of a petiole immediately after removal of the lamina will delay abscission, but that if the auxin application is delayed for some hours after lamina excision, then abscission may be accelerated rather than retarded. Because of this, it has been suggested that one can distinguish between phases in the abscission process—an initial “Stage 1” which is inhibited by auxin, and a “Stage 2” which may be promoted by auxin. It is likely that the explanation for the occurrence of these two stages is that during Stage 1 little or no senescence has occurred and applied auxin inhibits the senescence processes, but that by the time Stage 2 has been reached senescence is already occurring in the petiole of the explant and auxin is not able to bring it to a halt. As we have already considered, senescing tissues produce relatively large amounts of ethylene, and it is also known that auxin can promote ethylene synthesis. Thus, nodal explants in the senescent Stage 2 condition respond to added auxin by producing even more ethylene, and the abscission zone cells are already in an ethylene-sensitive condition, so that the result is an acceleration of the abscission process.

In summary, our present understanding of the regulation of leaf abscission envisages the following sequence of events: (a) the basipetal development of senescence of the lamina in response to an environmental stimulus or some internal “signal”, (b) the development of sensitivity to ethylene in the cells of the abscission zone, (c) a rise in ethylene levels in senescent cells, particularly those immediately distal to the abscission zone, to a critical level of approximately $1 \text{ nl g}^{-1} \text{ hr}^{-1}$, and (d) a series of biochemical and physiological responses to this ethylene in the cells of the abscission zone and in the nonsenescent cells lying immediately below the zone, which culminate in cell separation in the abscission layer. We consider below the cellular events which lead to cell separation, but before doing so it is appropriate to raise the question of what it is which controls the rate of production of ethylene in senescing cells distal to the abscission zone.

Evidence exists that during senescence of leaf cells there is release of a *non-volatile* substance which accelerates abscission. This has been termed the “senescence factor”, or “SF”, by Osborne. It has been suggested that abscisic acid (ABA) may be the SF, since applications of ABA can cause leaf senescence and abscission in a number of plant species. However, several lines of evidence argue against identifying SF as ABA. Thus, changes in endogenous ABA levels in leaves do not correlate well with time of leaf abscission, and it has been found that although exogenous ABA appears to affect directly the rate of ethylene production in some species, in others it appears to act by accelerating senescence directly but only indirectly inducing an increase in ethylene synthesis and rate of abscission. The SF, in contrast, is known to both accelerate abscission and have an *immediate* stimulatory effect on ethylene production. It would appear, therefore, that SF is a regulator of ethylene biosynthesis. The SF is present in both young healthy leaves as well as in old senescing leaves. However, SF is in a non-water soluble form in young leaves (but is extractable with organic solvents) but is water-soluble in senescing leaves. The significance of these observations on SF is not

at all clear at the present time, but it is tempting to speculate that the release of SF in leaf cells detected by Osborne may be the same phenomenon as the release of homocysteine from a cellular compartment in senescing flowers of *Opuntia tricolor* that results in increased ethylene synthesis (p. 66).

The act of leaf separation itself, which is the response induced by ethylene at an appropriate concentration, consists of two distinct processes: (a) enhanced cell growth in the region immediately proximal to (below) the separation layer, and (b) the synthesis and secretion of cell-wall-digesting enzymes by the same cells, which degrade the walls of the senescent cells above and lead to the separation of the two tissues. Growth of the cells proximal to the separation point is marked by rises in their rates of RNA and protein synthesis. A significant effect of this growth is to increase the physical pressure on the cells of the abscission layer, and also to provide the basis for the formation of a wound, or scar, tissue to seal the rupture surface after the leaf has been shed. The wall-degrading enzymes synthesized and secreted by the growing proximal cells consist of a range of appropriate polysaccharide hydrolases, including endopolygalacturonase and cellulase (β -1,4-glucanase), whose effects would be expected to be the weakening of cell to cell cohesion. The effect of ethylene on these enzymes is to enhance both their rates of synthesis and secretion through the plasmalemma of the proximal cells (Fig. 12.13). It is intriguing to note that the

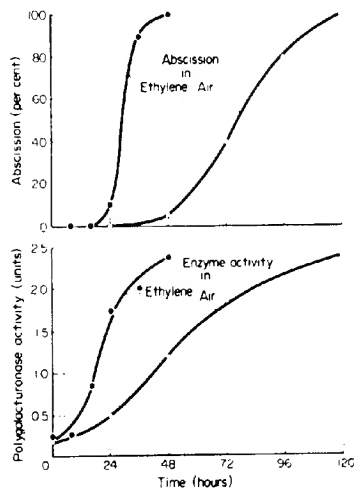


FIG. 12.13. A comparison of the time-courses of development of polygalacturonase activity and occurrence of abscission in *Citrus sinensis* leaf explants incubated in either air or in air containing 10 ppm ethylene. (Adapted from J. Rivov, *Plant Physiol.* **53**, 312-16, 1974.)

enzymes appear to be secreted only, or mainly, from one side of the layer of proximal cells; that which abuts the thin layer of abscission cells.

Once the walls of the abscission zone cells have been sufficiently weakened by the action of polysaccharide hydrolases, the final act of breakage is probably purely mechanical, aided not only by external forces such as wind action, but also by the cell enlargement on the proximal side of the separation layer and dehydration of the senescent tissues on the other, which together provide shearing forces in the abscission layer.

Fruit Abscission

Abscission phenomena are also shown by flowers and fruits. Thus, an abscission zone is commonly found at the base of the pedicel of the flowers in many species, if pollination and fertilization fail to take place. Similarly, even when successful fertilization has occurred, an abscission zone, leading to fruit-drop, may be formed at various stages during the development of the fruit. This is well seen in certain varieties of apple, in which there may be three peak periods of fruit drop: (1) immediately after pollination ("post-blossom" drop), (2) soon after growth of the young fruits ("June drop"), and (3) during ripening ("pre-harvest drop").

For some species it has been shown that the periods of fruit-drop coincide with periods of low auxin content in the fruit and conversely the times of low drop rate occur when the auxin content is high. This situation is found in the blackcurrant, in which we have already seen that it is possible to correlate growth rate with the levels of two auxins, one acidic and

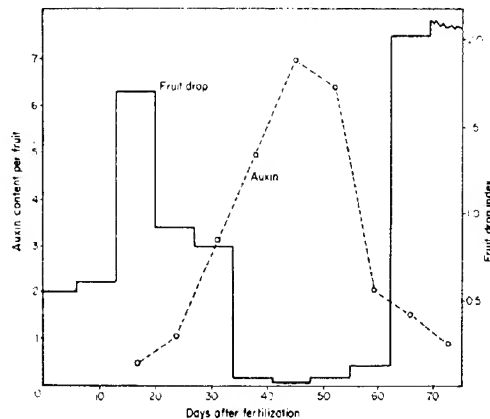


FIG. 12.14. Changes in the content of an unknown acidic auxin in the blackcurrant berry, in relation to the rate of fruit drop. (From S. T. C. Wright, *J. Hort. Sci.* **31**, 196, 1956.)

one neutral (Fig. 5.19). It has also been shown that there is a second acidic auxin the levels of which show a different pattern of variation which correlates inversely with the rate of fruit abscission (Fig. 12.14). This situation is analogous to that in leaves, therefore, where the formation of an abscission layer is associated with diminished auxin levels in the lamina.

On the other hand, in other species it has not been possible to correlate the variations in rate of fruit-drop with auxin levels and it seems likely that in such cases other factors play a role in determining fruit-drop. Indeed, there is little doubt that ethylene is involved in the control of fruit, as well as leaf, abscission, for this substance promotes fruit senescence and abscission.

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CHAPTER 13

The Control of Development

WE HAVE NOW considered plant development from several rather different standpoints. In the first two chapters we considered development from the viewpoint of the experimental anatomist and morphologist. Then we examined the action of growth hormones and discussed their roles in various aspects of growth and development. Finally we considered the physiology of flowering, senescence and dormancy, bringing in the effects of external factors, such as daylength, and internal factors, especially hormones. In the present chapter we shall examine some more general aspects of development.

GENES AND DEVELOPMENT

So far we have paid little attention to the genetical aspects of development, and yet every organism is, by definition, the product of the interaction between its genetic potentialities and the environment, and in the final analysis development has to be described in terms of activities of genes.

The fact that developmental processes are basically gene-controlled is self-evident, since genetic variations are known which affect almost every aspect of development, ranging from external morphology (leaf and fruit shape, flower colour, etc.) and internal anatomy to physiological characters, such as growth rate, flowering time and length of dormancy period. Thus, there seems no doubt that the pattern of development followed by any individual is determined primarily by the "programme" laid down in its genetic code.

During development there is progressive differentiation of organs and tissues, giving rise to a wide range of different types of cell. Not all the genes of the total gene complement are operative all the time and in all parts of the plant, however. Thus, the genes controlling flower development are apparently not normally operative in the embryo, nor during the purely vegetative phase of development. However, we know that the cells of a vegetative organ such as a leaf contain the genes required for flower development, since a complete new flower-bearing plant may be regenerated from leaf cells in certain species. Since, therefore, differentiation in plants apparently does not entail any genetic (i.e.

inheritable) differences between the nuclei of various types of cell and tissue, it must involve differences in *gene expression* in different parts and at different stages of the life cycle.

Since development is such an orderly process it must require that the right genes are expressed in the right cells at the right time. That is to say, development is essentially a process involving selective gene expression, and the concept that development involves the activity of specific groups of genes, which in turn control the synthesis of the enzymes and other proteins characteristic of specialized cells, is called the *variable gene expression theory* of differentiation. There must be some means of regulating the expression of specific genes during cell differentiation and, since development is a very orderly process, there must be some mechanism for determining the sequence of gene expression. Therefore we have to consider how selective gene expression may be achieved.

The Control of Gene Activity

Gene expression involves (1) transcription of the DNA sequences to form messenger RNA, (2) translation of the information in the m-RNA into amino acid sequences in the polypeptide chain of enzyme and structural proteins, (3) activity of the enzymes in various reactions to give *gene products*. The essential features of these processes are too well known to require detailed elaboration here. Before we can discuss the control of selective gene expression in development, however, we need to consider how *gene activity*, at the transcriptional level, may be regulated. Information on this subject is almost entirely restricted to prokaryotic micro-organisms, especially the bacteria, which we will consider first.

Bacteria

Some enzymes of bacteria are present all the time and are said to be constitutive. On the other hand, other enzymes are only formed when their substrate is present in the external medium. For example, when the bacterium, *Escherichia coli*, is grown in the absence of a galactoside (i.e. a compound containing the sugar, galactose, linked to another, non-sugar molecule), only traces of the enzyme β -galactosidase are formed, but as soon as a galactoside is added, the rate of synthesis of this enzyme increases enormously. This type of enzyme is said to be *inducible*. Removal of the substrate results in the almost immediate cessation of enzyme synthesis.

Contrasting with this process of enzyme induction is the *repression* of enzyme synthesis seen in other enzyme systems. For example, when *E. coli* is grown in the absence of the amino acid histidine, the enzymes involved in the production of histidine are actively synthesized. As soon as histidine is added to the medium the enzymes cease to be synthesized. In this instance, therefore, there is *repression* of enzyme synthesis. This phenomenon is called "end-product repression", since the product of the sequence of reactions, in this case histidine, inhibits the formation of the enzymes concerned with its own biosynthesis.

The result is that if the end-product is supplied from outside, it inhibits its own synthesis, and hence the cell ceases to make any more of the compound.

The phenomenon of enzyme induction and repression led Jacob and Monod to propose a general theory for the control of gene activity in bacteria, which we shall now describe briefly. The gene which determines the structure of a specific enzyme protein is referred to as a *structural gene*. The synthesis of m-RNA is assumed to be initiated only at certain regions or *operators* of the DNA strands (Fig. 13.1). In some instances a single operator may control the transcription of several adjacent structural genes into m-RNA. The segment of DNA thus controlled by a single operator is referred to as an "operon", which may contain one or several structural genes.

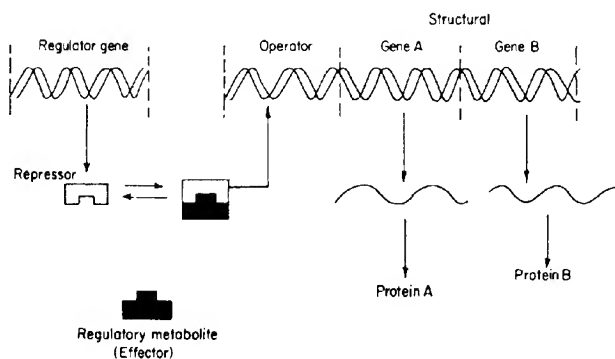
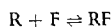


FIG. 13.1. Scheme illustrating the control of protein synthesis according to the theory of Jacob and Monod (see text).

The rate of transcription of structural genes is controlled by other genes referred to as *regulator genes*. A regulator gene forms a cytoplasmic product or *repressor*. The repressor formed by a given regulator gene is assumed to have an affinity for certain specific operator genes, with which it binds. This combination blocks the production of m-RNA by the whole operon controlled by the operator, and therefore prevents the synthesis of the specific proteins controlled by the structural genes of the operon.

A repressor (R) has the property of reacting with certain small molecules, called "effectors" (F) expressed as follows:



In inducible systems only the R form of the repressor is active and blocks the transcription of the operon. The presence of an effector (or inducer) inactivates the repressor and therefore allows messenger-RNA synthesis to proceed. By contrast, in repressible systems only the combined RF form of the repressor is active. Synthesis of m-RNA by the operon, allowed

in the absence of the effector (or repressing metabolite), is, therefore, prevented in its presence. These ideas are summarized in Fig. 13.1.

The repressor may be an "allosteric" protein, with two active sites, one able to react with the operator and one to react with the inducing or repressing molecule. It is supposed that when the repressor combines with an inducer its shape is deformed somewhat, so that it can no longer react with the operator, which is thereby "derepressed".

There is much evidence in support of the existence in bacteria of the type of regulator system postulated by Jacob and Monod, but there is no evidence to suggest that it exists also in higher plants. Moreover, even in bacteria the Jacob and Monod hypothesis apparently applies to the regulation of only a few enzymes. However, genes are known in maize which appear to show some of the characters of regulator genes. Thus, a locus in maize known as "Activator" (Ac), appears to be a master locus for a second locus, "Dissociation" (Ds), which is unable to function in the absence of Ac. Ds, in turn, affects the expression of a number of other genes, and hence is analogous to an "operator" gene in bacteria, while Ac may be regarded as the "regulator".

For example, under certain conditions Ds causes C, which gives coloured aleurone in the grains, to behave as if it were the colourless recessive allele, *c*. Thus Ds appears to "repress" the action of C. On the basis of these and other observations, McClintock has postulated that the chromosomes of higher plants contain both genes and "controllers", which regulate the action of the genes.

It is not difficult to construct models, based upon the Jacob and Monod theory, which would explain several features of differentiation in plants, as we shall see later (p. 313).

Higher Plants

We do not yet know whether the bacterial type of control system is operative in eukaryotic organisms, including higher plants, in which the DNA is contained in a nucleus and organized into chromosomes at nuclear division. In fact, evidence is accumulating which suggests that a somewhat different mechanism may be involved in the control of gene activity in higher plants and animals.

It has long been known that the nuclei of plant and animal cells contain not only DNA, but also significant amounts of protein, most of which is histone, a type of protein which is basic in nature, due to the high content of the basic amino acids, lysine and arginine. The possible role of histones in repressing DNA has been studied, particularly by J. Bonner and his co-workers, by using preparations of chromosomal material referred to as "chromatin". Chromatin is prepared by disruption of the tissues by homogenization, filtering and selective sedimentation by centrifugation. In this way relatively pure interphase chromosomal material may be prepared and its properties studied *in vitro*. Such chromatin contains DNA, histone, a small amount of non-histone protein and a small amount of RNA. It may also contain the enzyme RNA-polymerase, which is responsible for the linking of nucleotides during the synthesis of m-RNA.

Isolated chromatin is capable of synthesizing RNA when supplied with the nucleoside triphosphates of the four RNA bases, guanine, adenine, cytosine and uracil. It is thus possible to study the RNA-synthesizing capacity of DNA from various types of plant material.

It has been shown, for example, that chromatin extracted from dormant potato buds has lower RNA-synthesizing ability than chromatin from non-dormant buds, suggesting that possibly dormancy involves rather extensive repression of DNA. Similar results have been obtained with chromatin prepared from dormant and non-dormant hazel seeds. These observations do not, of course, show that histone is involved in the repression mechanism, the chief evidence for which is provided by the finding that when the histone is removed from chromatin, the deproteinized DNA is found to have very much greater RNA-synthesizing activity than chromatin not so treated. This has been reportedly demonstrated by the finding that removal of the protein, including the histone, from pea-bud chromatin derepresses the genes for globulin synthesis.

One difficulty for the hypothesis that histones act as gene-repressors is the fact that relatively few types of histone are known to occur naturally, whereas there is a large number of genes, each of which would seem to require a specific repressor. A further problem is the fact that histones are not present in the bacterial chromosome and yet there is effective gene activation and repression in bacteria. Indeed, the repressors in bacteria appear to be specific, non-histone proteins. It seems likely, therefore, that if histones do function as gene repressors they can only do so in relatively non-specific manner, as possibly in dormancy, and that the fine control of gene activity is effected by non-histone proteins. Indeed there is evidence that regulation of specific gene activity may involve acidic proteins of which a considerable number of different types are present in the nucleus. Furthermore, work on plants suggests that acidic nuclear proteins may regulate RNA synthesis and the course of differentiation.

Levels of Control of Gene Expression in Development

Control of selective gene expression in development has been postulated to be effected at various stages in the chain of processes occurring between the gene and its end product, including the following:

- (1) *transcription*, i.e. by the selective transcription of specific genes;
- (2) *translation*, i.e. by selective translation of specific m-RNAs at different stages of differentiation;
- (3) *enzyme activation*, i.e. by selective activation of pre-existing enzymes.

The most direct and efficient method of controlling selective gene expression would appear to be at the transcription stage, since most messenger RNAs appear to have a rather short life and it would appear to be inefficient if a large number of m-RNAs are continually produced, only a few of which are engaged in protein synthesis at any one time.

It seems very probable, on *a priori* grounds, therefore, that development involves the selective activation and repression of specific genes in the proper sequence, but since we

do not know how gene activation is controlled in eukaryotes, including higher plants, and we have no means of separating and identifying specific m-RNAs, this hypothesis cannot be tested by direct experiment at present. However, by using inhibitors of transcription, such as actinomycin D, it is possible to demonstrate indirectly that various aspects of development, such as flowering (p. 243), are probably dependent upon the synthesis of new m-RNAs.

On the other hand, there are instances where control appears to be effected at the translation level in plants, as in the germination of seeds (p. 280). There is also increasing evidence for the existence of mechanisms for controlling selective enzyme activation.

It is clear that control of translation and enzyme activation must occur in the cytoplasm, whereas transcriptional control must occur in the nucleus. However, even events occurring in the nucleus are profoundly influenced by the cytoplasmic environment in which it is sited. Indeed, there is a complex interplay between the nucleus and the cytoplasm which is well illustrated by studies on the marine alga, *Acetabularia*, and the aquatic fungus, *Blastocladiella*.

Acetabularia consists of a single, giant cell, which has a rhizoid and a cylindrical stalk (Fig. 13.2). The stalk grows in length at its apical end and ultimately forms a circular cap

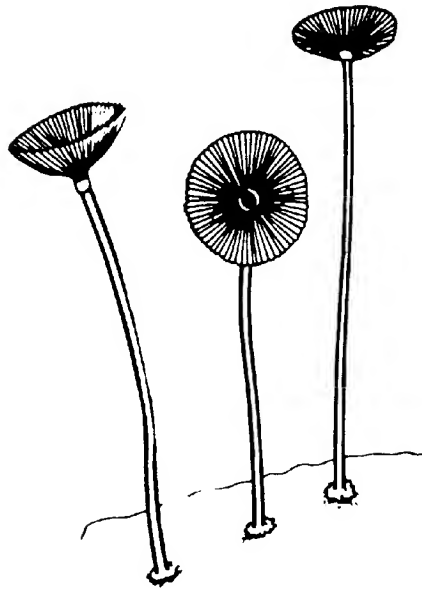


FIG. 13.2. Plants of *Acetabularia mediterranea*.

consisting of a considerable number of sections called "rays", in which large numbers of cysts are ultimately formed and released; the cysts give rise to isogametes after shedding. There is a large central vacuole, with a boundary layer of cytoplasm inside the wall, and the single nucleus lies in the cytoplasm at the rhizoid end.

Acetabularia shows considerable capacity for regeneration and if the stalk is cut off above the rhizoid the latter will regenerate a new stalk and cap. However, even parts lacking a nucleus can regenerate. For example, if a young stalk which has not yet developed a cap is separated from the rhizoid, so that it lacks a nucleus, it will survive for a long time and will ultimately form a cap.

Different parts of the cell may be grafted together. If the apical part of the young stalk is cut off and pushed over the basal part of another plant containing a nucleus, it will continue to grow and develop normally. Grafting may also be carried out between various species of *Acetabularia*, which differ in the shape of the cap. If an anucleate section of one species is grafted to a nucleate base of another species, the regenerated plants show mainly the characters of the species which supplied the nucleus.

These various results seem to show that development is ultimately under the control of the nucleus, but that there are long-lasting effects of the nucleus on the cytoplasm, which can continue and complete morphogenesis even if the nucleus is removed. It has been suggested that certain "morphogenetic substances", possibly messenger-RNA formed in the nucleus, are present in the cytoplasm and persist over long periods.

Although cap formation depends upon the presence of stable morphogenetic substances formed in the nucleus, the expression of the genetic information is controlled and effected by regulatory mechanisms in the cytoplasm. The information passed from the nucleus to the cytoplasm can evidently remain unexpressed for several weeks, and the time of cap formation appears to be determined by events that take place in the cytoplasm rather than the nucleus. Control of gene expression in *Acetabularia* therefore appears to be at the translational level.

These latter conclusions are supported by the results of studies on the aquatic fungus, *Blaschkeella emersonii*. The zoospores of this species, which lack cell walls, have a single, posterior flagellum. There is a nuclear cap which surrounds one end of the nucleus and contains large numbers of pre-formed ribosomes. The zoospores are released into the water and settle on a solid substratum where they enter a short period of encystment, during which they undergo rapid and radical changes in structure. These changes include the breakdown of the nuclear cap and release of the ribosomes, which become dispersed throughout the cell. In about 10 minutes a small germ tube appears, indicating the commencement of germination (Fig. 13.3).

Studies on protein and nucleic acid synthesis have shown that RNA synthesis does not begin until much later (40-45 minutes after encystment) and this is followed in a further 30-40 minutes by protein synthesis and then DNA synthesis. Thus, germination takes place apparently without any RNA or protein synthesis. Moreover, inhibitors of RNA and protein synthesis do not prevent encystment and germination, so that it would appear that the necessary ribosomal transfer and messenger RNAs are all present in the zoospores

before they encyst, and polysome formation can be seen to occur during early germination. These structural changes occurring during encystment and germination apparently require only the protein and RNA present in the zoospore when it is released, but the later stages of germination do appear to require new protein synthesis.

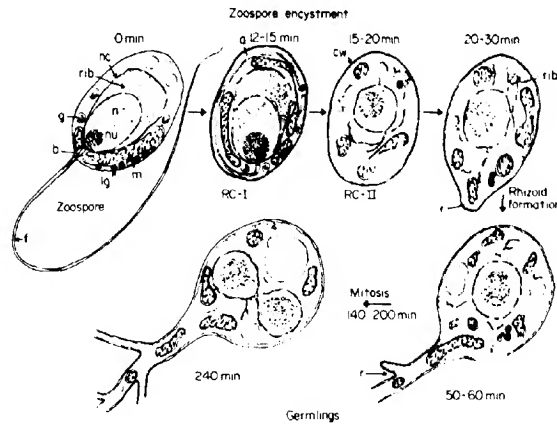


FIG. 13.3. Diagram of zoospore germination and early development in *Blastocladiella emersonii*. Symbols: b, basal body; g, gamma particle; m, mitochondrion; rib, ribosomes; nc, nuclear cap; nu, nucleolus; n, nucleus; lg, lipid granules; a, flagellar axoneme; cw, cell wall; r, rhizoid; f, flagellus; v, vacuole; RC-I, round cell I; RC-II, round cell II. (From C. J. Leaver and J. S. Lovett, *Perspectives in Experimental Biology*, Ed. N. Sunderland, Pergamon Press, Oxford, pp. 229-311, 1976.)

We have also seen that there is evidence that protein synthesis during the early stages of germination of seeds depends upon pre-formed m-RNA (p. 280). Thus, although it seems very probable that development involves selective gene activation and repression at the DNA level, the m-RNAs released into the cytoplasm are not necessarily immediately involved in protein synthesis, and the translation process offers another stage at the time of gene expression may be controlled. Other types of control of gene expression have been suggested but there is little conclusive information on this crucial question for an understanding of development.

Biochemical Differentiation in Higher Plants

Ultimately the processes occurring during cell differentiation are terminated and the cell reaches the "steady state" of the mature condition, in which metabolism is maintained continuously (except, of course, in the case of non-living cells such as those of the xylem). The visible manifestations of differentiation include variations in the development of the cell wall and of certain cytoplasmic organelles, such as plastids. It is clear that differentiation must also extend to certain aspects of metabolism when it is remembered that some tissues are specially adapted for particular functions, such as photosynthesis, secretion and storage of reserve materials. Such differentiation almost certainly involves differences in enzyme production, which in turn implies the maintenance of differences in gene activation and repression between various cells even in the mature state.

Many basic metabolic pathways are probably operative in all living cells of the plant. This must apply to the main pathways involved in the respiratory breakdown of carbohydrates, for example. On the other hand, there is much evidence that various tissues differ in their biosynthetic abilities. For example, it is found that the isolated roots of many species require the supply of certain vitamins, including thiamin, pyridoxine and nicotinic acid when grown in aseptic culture. It appears that in the intact plant these vitamins are synthesized in the shoot and are supplied to the roots. Similarly, callus cultures of certain plant tissues, including tobacco pith tissue, require to be supplied with auxin and cytokinin in order to maintain cell division in culture (p. 145). Evidently tobacco pith cells do not have the ability to synthesize auxin and cytokinin and hence require an exogenous supply. However, this inability of pith cells to synthesize the two hormones is not due to any permanent loss of the potentiality to do so, as shown by the fact that cytokinin-requiring callus cultures can be "habituated" (p. 145), so that they become cytokinin-independent. Studies on the bacterial disease of plants, known as "crown gall", also have a bearing on this topic. Infection by *Agrobacterium tumefaciens* brings about the transformation of normal plant tissue into tumour tissue, and once transformed it will continue to grow as tumour tissue indefinitely (p. 145). As a result of infection the normal tissue has apparently undergone a profound and stable change. It is found that the tumour tissue will grow actively in sterile culture without the addition of auxin and cytokinins, indicating that it can now synthesize its own hormones. Transformation is brought about by the transfer of a section of bacterial DNA ("plasmid") to the plant cell. Presumably this bacterial DNA segment either carries genes for cytokinin biosynthesis or causes activation of the section of the plant genome controlling cytokinin biosynthesis.

The inability of roots to synthesize certain vitamins, and of tobacco stem tissue to synthesize auxins and cytokinins, provides rather strong evidence that cell differentiation involves the activation of certain genes and the repression of others. It would be of interest to know whether the meristematic cells of the shoot apex of tobacco have the ability to synthesize cytokinins; if so, then it would seem that one of the processes occurring during cell differentiation in the stem is the repression of the enzymes responsible for auxin and cytokinin synthesis. Such a change in synthetic ability might, indeed, explain the transition

from cell division to cell expansion seen in the apical region of both shoot and root. On the other hand, all cells of the shoot may lack the capacity for biosynthesis of cytokinins and may depend upon supply of this hormone from the roots (p. 290).

We do not know how these permanent differences in biosynthetic ability are maintained but Jacob and Monod have shown that it is not difficult to construct "model" circuits, based upon the bacterial repression concepts, which would result in certain genes being permanently switched on. One very simple circuit is shown in Fig. 13.4, in which the

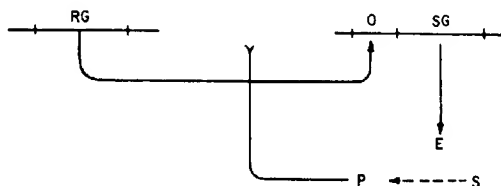


FIG. 13.4. Hypothetical model of a circuit for control of enzyme synthesis in which the inducing substance is the product of the controlled enzyme. Synthesis of enzyme E, genetically determined by the structural gene SG, is blocked by the repressor synthesized by the regulator gene R.G. The product P of the reaction catalysed by enzyme E acts as an inducer of the system by inactivating the repressor, O, operator; S, substrate. (From F. Jacob and J. Monod, *Cyto-differentiation and Molecular Synthesis* (Ed. M. Locke), Academic Press, New York and London, 1963.)

inducer is not the substrate but the product of the controlled enzyme system. Such a system is known to occur in bacteria. In the absence of an exogenous inducer, the enzyme will not be produced, unless already present, but temporary contact with an inducer will cause the system to become "locked" indefinitely, at least so long as either substrate or product is present. Suppose, for example, that the operon in Fig. 13.4 is responsible for the biosynthesis of a hormone, such as cytokinin or gibberellin, and that in a dormant seed the operon is repressed. Then a single treatment with exogenous hormone will activate the operon and from then on the seedling would be able to synthesize its own hormone. The stimulation of endogenous auxin synthesis by a single application of exogenous IAA in fruit set (p. 125) may provide a similar example. It is quite easy to construct other models which would account for the permanent suppression of certain enzymes. Thus, in the model illustrated in Fig. 13.5, the product of one enzyme acts as an inducer for the other, so that the two enzymes are mutually dependent. One could not be synthesized in the absence of the other, and inhibition of one enzyme or elimination of its substrate, even temporarily, would result in the permanent suppression of both.

There is no evidence that determination and differentiation involve this type of mechanism, but these circuits do illustrate that it is possible to construct models which would account for the observed facts, within the concepts of bacterial repressor mechanisms.

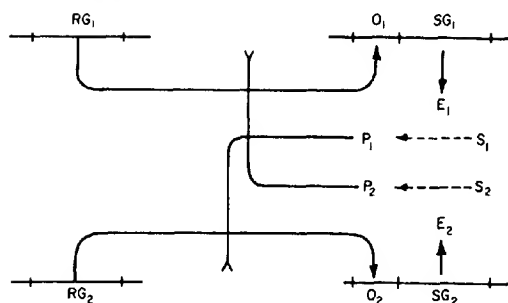


FIG. 13.5. Model circuit in which the product of one enzyme acts as an inducer of the other. Synthesis of enzyme E_1 , genetically determined by the structural gene SG_1 , is blocked by the repressor synthesized by the regulator gene RG_1 . Synthesis of another enzyme E_2 , controlled by structural gene SG_2 , is blocked by another repressor, synthesized by regulator gene RG_2 . The product P_1 of the reaction catalysed by enzyme E_1 acts as an inducer for the synthesis of enzyme E_2 , and the product P_2 of the reaction catalysed by enzyme E_2 acts as an inducer for the synthesis of enzyme E_1 . O, operator; S_1 , S_2 , substrates. (From F. Jacob and J. Monod, *Cytodifferentiation and Molecular Synthesis* (Ed. M. Locke), Academic Press, New York and London, 1963.)

THE CONTROL OF DIFFERENTIATION

The Canalization of Development

So far, in this chapter, we have been considering some of the possible mechanisms which control the activation and repression of genes. We now have to consider these processes in relation to development of the organism as a whole.

We have seen that development of a flowering plant involves a series of successive stages, starting with the differentiation of the embryo into root and shoot, followed by the formation of organ initials, and ending with tissue differentiation within the individual organs. Superimposed on this pattern are the major changes in development represented by the transition to the flowering phase and the onset of dormancy. The orderly manner in which the successive stages follow each other in a regular sequence is one of the most striking features of development. This phenomenon is well illustrated in the development of a flower, where not only is there a regular sequence in the initiation of the various flower parts (perianth, stamens and carpels), but within each of these there is an equally regular pattern of development.

The transition from one stage of development to the next would seem to involve a process of successive gene activation, in which certain previously repressed genes come into action and others become or remain repressed. Direct evidence of successive gene activation is provided by insect development. As is well known, the cells in some organs of *Drosophila* and certain other flies have "giant" chromosomes, formed by the repeated replication of the DNA strands, without nuclear division. These giant chromosomes thus each consists of large numbers of single chromosome threads lying parallel to each other and

aligned so that corresponding regions of the various threads are opposite to each other, to give a characteristic banded appearance. Each band appears to correspond to a single gene or operon. At certain stages of development of the insect one or more of these bands swells up and forms a "puff" of what is apparently RNA. It would seem that the occurrence of puffing indicates that a particular gene is active at that time. Different tissues have characteristic puffing patterns, and these occur at specific periods of development. Moreover, if the moulting hormone, ecdysone, is administered to a larva the puffing pattern changes rapidly, as the insect enters a new phase of development. We have no direct evidence of successive gene activation during development in plants, but it seems very likely on *a priori* grounds, that this does occur.

If development involves an orderly sequence of expression of large numbers of genes, this raises the question as to how such gene control is achieved. Clearly specific genes must be expressed in the right cells at the right time.

The successive stages of development can be regarded as a process of "programme selection", in which there is divergence into alternative pathways of further development at various critical points of time and space. This divergence may occur at the cellular level, as when two daughter cells of a common mother cell show divergent patterns of differentiation, or it may occur in the differentiation of organs, or even in the shoot apex as a whole, as in the transition from the vegetative to the flowering phase. Furthermore, we saw that once an organ, such as a leaf primordium, passes a certain stage of development, it becomes irreversibly "determined" as a leaf (as opposed to a bud) and cannot normally then be converted into any other structure (p. 35).

This successive commitment of different parts of the developing organism to specific pathways of differentiation has been referred to as the "canalization of development". The development of the embryo appears to involve canalization into several major pathways, as a result of which there is determination of root and shoot regions, and with the establishment of an organized shoot apex the basis is laid for the initiation of various organs (leaf, bud, stem), which is maintained throughout the subsequent vegetative phase of the plant.

It will be seen that the general features of development raise a number of major problems viz. (1) how are divergent patterns of differentiation initiated in cells which previously shared a common origin?; (2) how are such differences maintained through subsequent cell divisions?; (3) how is the regular sequence of successive changes seen in development achieved?

Our knowledge of these matters is still very fragmentary, but we shall discuss the first two problems in relation to the phenomena of *unequal cell division* and *determination*. We have no knowledge of how the co-ordinated sequence of events in development are achieved but the regular sequence of changes seen in the development of an organ such as a leaf or a flower strongly suggests that once these particular pathways have been entered, then all the subsequent stages follow inexorably, as "chain reactions" in which the attainment of one stage seems to trigger off the next one. If this latter type of mechanism is involved, then it would imply that the pattern of developmental events is predetermined, at least for

certain organs, and that development is primarily controlled by an internal mechanism, as if the organism is an internally programmed piece of equipment which, once it is switched on, will automatically go through a regular sequence of activities, each of which is triggered off by the preceding one.

A mechanism of this type has been suggested for flower development. It is postulated that once the transition to the flowering stage has begun, a gene complex A is activated in the first formed primordia, and these genes produce an inducer X which moves to the next set of primordia and there activates a gene complex B, and so on through the different classes of organ. This hypothesis postulates the occurrence of short-range intercellular "messengers" or "hormones", but the existence of such substances has yet to be demonstrated.

By contrast with the "internally programmed" type of development we have just been considering, we have already seen that certain stages of the life cycle, notably flowering and bud dormancy, may be controlled by environmental factors such as daylength and temperature. In plants showing photoperiodic control of flowering, it is clear that the onset of flowering is not rigidly predetermined as part of a sequence of internally programmed changes, but may be controlled by a factor of the external environment. In this case the daylength conditions may control gene switching through phytochrome.

Thus, we apparently have two rather contrasting types of control mechanism. Internal programming, such as we see in leaf and flower development, appears to be particularly characteristic of organs of determinate growth, whereas environmental control appears to be more common with changes involving the shoot apex, as in the initiation of flowers and dormant resting buds. However, this distinction between the development of organs of determinate and indeterminate growth patterns is not absolute, since light also plays an important part in normal leaf development (p. 196).

Unequal Cell Division

One of the central problems of development concerns the nature of the processes whereby cells of common lineage are caused to diverge into alternative pathways of differentiation. One such process is by *unequal division* of a parent cell, leading to two daughter cells which subsequently follow divergent patterns of differentiation.

This process can occur at various stages in the life cycle, from the initial division of the zygote to the final stages of cell differentiation in an organ such as a leaf.

As we have seen (p. 24) the early development of the zygotes of flowering plants also involves an unequal division but the orientation of the spindle of the first division is evidently determined by the surrounding maternal tissues of the ovule, so that the root end is always towards the micropyle and the shoot end away from it. The cytoplasm of the unfertilized egg of *Capsella* is highly polarized for up to one-half of the micropylar end is filled with a large vacuole, whereas the chalazal end contains the nucleus and much of the

cytoplasm (Fig. 13.6). The first division gives rise to a smaller, densely cytoplasmic cell which forms most of the future embryo, and a larger vacuolated cell which forms the suspensor.

Unequal division also plays an important role in cell differentiation in the later stages of development of higher plants. In the root epidermis of some species, including certain grasses, root hairs arise from daughter cells formed by the unequal division of certain epidermal cells. These cells have their long axis parallel to the root axis, and it can be seen that the cytoplasm of these cells is more dense at the apical end. Mitosis occurs and a transverse cell wall is formed in a position which gives rise to a small daughter cell with dense cytoplasm and a larger cell with less dense cytoplasm (Fig. 13.7). The root hair initials are normally formed only from the smaller cells.



FIG. 13.6. Electron micrograph of the zygote of *Capsella bursa-pastoris*, showing polarization of the cell as manifested by the occurrence of the nucleus lying in the dense cytoplasm at the micropylar end (left) and a large vacuole at the chalazal end (right). (From Sister Richardis Schultz and W. A. Jensen, *Amer. J. Bot.* 55, 807-19, 1968.)

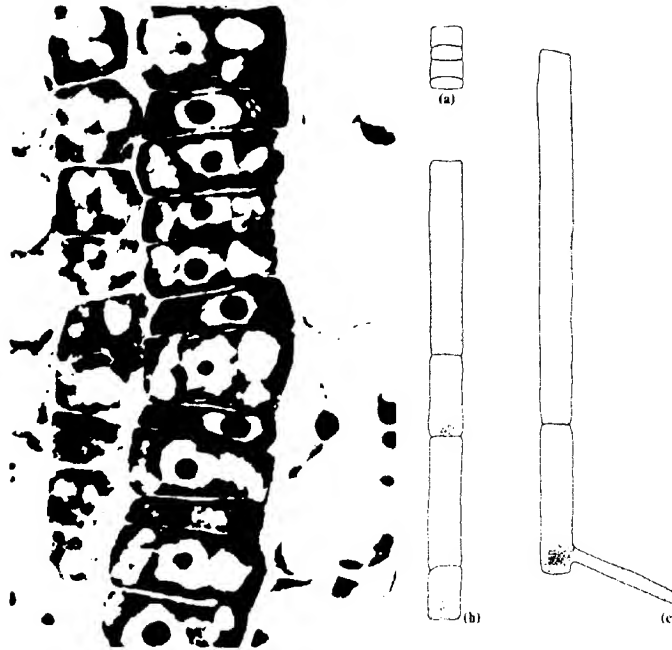


FIG. 13.7. *Left*: Formation of trichoblasts (root hair initials) in the root epidermis of *Hydrocharis morsus-ranae*. Unequal division has given rise in each case to a small cell which is the trichoblast and a large cell which is the epidermal cell (c). (From E. G. Cutter and L. J. Feldman, *Amer. J. Bot.* 57, 190-201, 1970.)

Right: Development of root hair initials in the grass *Phleum pratense*, at successive stages of development (a-c). The smaller cells formed by unequal division (a) gives rise to the root hair cell (c). (From E. W. Sinnott and R. Bloch, *Proc. Nat. Acad. Sci., U.S.A.* 26, 223-7, 1939.)

A similar situation is seen in the development of the stomatal guard cells of certain monocotyledons. Here also, certain epidermal cells of the developing leaf show unequal division, and cut off a small cell with densely staining cytoplasm at one end, and a larger cell at the other (Fig. 13.8). The smaller cell becomes a stomatal mother cell and it undergoes a further division, at right angles to the first division, and the two daughter cells so formed are in this case identical and give rise to the guard cells. Thus, where a parent cell shows polarized differences in the cytoplasm along its axis, division is unequal and leads to differentiation between the resulting daughter cells, but where the plane of division appears to divide the cytoplasm equally, the resulting daughter cells are identical.

The fact that unequal division is preceded by polarized differences in the cytoplasm is

well illustrated in the development of pollen grains in which the spindle is oriented so that one of the daughter nuclei passes to the end of the cell at which there is denser cytoplasm, and becomes the generative nucleus, whereas the other daughter nucleus moves to the region of the cell with less dense cytoplasm and becomes the vegetative nucleus (Fig. 13.8B). Occasionally the plane of the spindle accidentally becomes oriented *across* the axis of the pollen mother cell, and in this case two equal cells are formed and the further development of the pollen grain is disturbed (Fig. 13.8C).

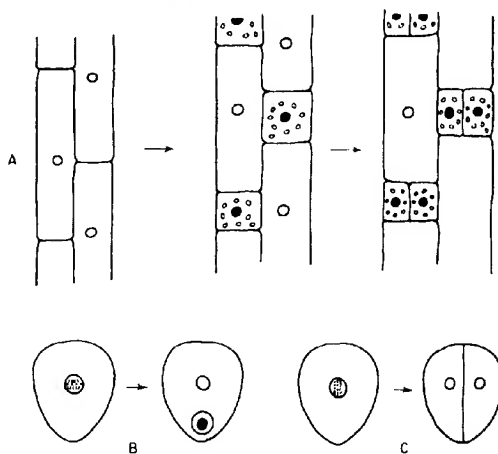


FIG. 13.8. A. Unequal division in formation of stomatal guard cells in leaf of a monocotyledonous plant. Epidermal cells undergo unequal division with the formation of a smaller stomatal mother cell (nucleus shaded) and a larger cell (nucleus not shaded). Chloroplasts develop in the stomatal mother cell, which divides again at right angles to the plane of the first division to produce two daughter cells which develop into the guard cells.

B, C. Development of pollen grain.

B. Normal development. Nucleus divides so that one of the daughter nuclei moves to the region of more dense cytoplasm at one end of the cell and becomes one generative nucleus. The other daughter cell nucleus becomes the vegetative nucleus.

C. Abnormal development. Nucleus divides at right angles to the normal plane. Daughter nuclei thus remains in the same cytoplasmic environment, and two equal cells are formed which disrupts the further normal development of the pollen grain. (Redrawn from E. Bünning, *Handbuch Protoplasmaforschung*, Vol. VII, Vienna, 1958.)

It is not known how the polarized differences in the cytoplasm in the mother cell arise in the first instance, but the process of division itself is likely to lead to polarization, since the end of a daughter cell at which the equator was formed is likely to contain a different distribution of organelles from the end at which the pole of the spindle occurred.

In most of the examples we have just considered, the cells derived from unequal division

do not themselves undergo further division, but differentiate directly. In some instances, however, the derivative cells from an unequal division undergo several more divisions. For example, in plants of the castor bean (*Ricinus communis*) there are secretory cells which contain tannin and unsaturated fatty acids. These cells rise from an initial unequal division, and one daughter cell undergoes a series of divisions to give rise to a row of cells, each of which becomes a secretory cell. Thus, the differential state can apparently be transmitted by cell lineage in some instances.

The question arises as to whether unequal division, followed by transmission of the resulting differences through further cell division (i.e. by cell lineage), are normally involved in other aspects of differentiation. When we consider certain lower plants, especially the algae and bryophytes, the importance of unequal division in differentiation is quite clear. For example, in the alga, *Chara*, and in mosses and many ferns, the single apical cell divides unequally, the outer daughter cell continuing as the apical cell and the other giving rise to differential thallus cells after a second unequal division (Fig. 13.9). In certain mosses and a few pteridophytes, such as certain species of *Selaginella*, the various tissues of the mature stem can apparently be traced back to precise divisions of the single apical cell and its immediate derivative cells, so that differentiation here also appears to arise by unequal division followed by transmission by cell lineage. However, in the shoots of other pteridophytes and of seed plants, it is not possible to trace any clear relation between the pattern of cell division at the shoot apex and the differentiation of the tissues derived from it.

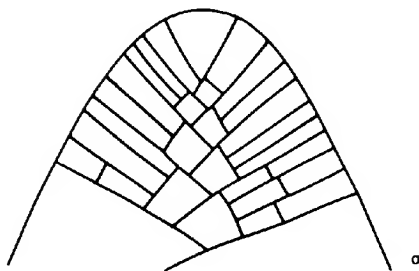


FIG. 13.9. Segmentation of a tetrahedral apical cell of a lower plant as seen in median section. (From Clowes, 1961.)

DETERMINATION IN PLANT CELLS

We have been considering the various processes whereby cells which hitherto had showed a common lineage diverge into alternative pathways of development. Once a group of cells has become committed to a particular pathway it usually follows the "normal" pattern of development through to completion and it is unusual for it to revert to an earlier stage or to make the transition into some other pathway. Thus, leaf primordia

do not become buds or stems, and although abnormalities may sometimes occur during flower development, with reversion to a vegetative apex, such instances are relatively rare, and at certain critical stages a particular part of the organism is said to become "determined" in respect of its further differentiation. We have already seen an example of such determination in the development of leaf primordia (p. 37). How this irreversible commitment which occurs during determination is brought about is completely unknown, but it seems necessary to make a distinction between (1) a situation in which the pattern of development is controlled by influences *outside* the cell and (2) one in which the cells undergo changes which result in *intrinsic* differences in their developmental potentialities. An example of the first situation is seen in the transformation of an apical meristem from the vegetative to the flowering condition in response to a photoperiodic stimulus. As we have seen, hormonal factors appear to play an important role in this developmental switch. The second situation, in which intrinsic differences arise during development has been little studied in plants, although its occurrence in animals is well established, as shown by the various phenomena included under the heading of "cell determination", in which the possible paths of differentiation of a given cell lineage become progressively restricted during the development of an animal embryo. Probably the reason for this neglect of the phenomena by botanists is that they have been fascinated by the "totipotency" of plant cells, as illustrated by the capacity of differentiated cells, such as leaf cells, to regenerate whole plants and the potentiality for producing embryos from almost any part of the carrot plant (p. 149). However, evidence for the occurrence of determination in plant cells and tissues is not lacking and it is shown most clearly in shoot apices.

It might seem paradoxical to suggest that meristems exhibit determination, since meristematic cells are normally regarded as essentially cells which are undifferentiated and uncommitted. However, we have already considered two or three examples of determination in shoot apical meristems, viz. in vernalization and in phase change in woody plants. As we have seen, vernalization of rye embryos leads to intrinsic changes in the developmental potentialities of the cells of the shoot meristem so that they become capable of flower initiation. Once the apical meristem of rye has become fully vernalized all the cells derived from it are in the vernalized state, which is apparently transmitted through cell division without "dilution" (p. 232). Although the vernalized state is a very stable one, the new embryos produced by a vernalized plant are not themselves vernalized and hence "devernalization" must occur during normal gametogenesis.

"Phase-change" in woody plants (p. 250) provides another example of determination in apical meristems. There is no doubt that phase change involves intrinsic changes in meristematic cells, since if sterile callus cultures, derived from juvenile and adult stem tissue of ivy, are grown on the same nutrient medium, differences in the two cultures can be recognized, with "juvenile" cultures maintaining a higher rate of growth and more abundant root regeneration than the "adult" cultures. Evidently we are dealing with two relatively stable alternative states of determination which do not involve genetic change (since the adult shoots produce seeds from which juvenile seedlings develop), but which can be transmitted through cell division without loss of determination.

A further example of stable differences in shoot apical meristems is provided by the alternation of free living gametophytes and sporophytes in ferns and other lower plants. Normally the gametophyte is haploid and the sporophyte is diploid, but the differences in morphology and life cycle between the two generations cannot be ascribed to the difference in ploidy, since it is possible to produce experimentally diploid or tetraploid gametophytes and haploid sporophytes.

These various phenomena must all depend upon activity of different parts of the genome in the alternative states. The differences between shoot and root apices may also provide an example of determination, since they are stable structures which normally retain their identity over many cell generations.

Differentiation in Apical Meristems

We have so far considered from a general standpoint the concepts of unequal division and determination. We now have to consider how far we can apply these concepts to interpret differentiation at various levels, viz. as between shoot and root, and at the organ, tissue and cell levels, as established at the apical meristems.

We have seen that the establishment of root and shoot poles in the embryo of higher plants is preceded by an unequal division in the zygote. Thus, the major division of the plant body into root and shoot is established at the earliest stages in development. Once established, the shoot and root meristems are very stable structures, and it is extremely rare for a shoot meristem to be converted into a root meristem and vice versa. Moreover, the stability of the root meristems is intrinsic in those organs themselves and is not dependent on influences from other parts of the plant, since isolated roots of tomato and other species can be maintained in aseptic culture for many years without the appearance of any buds, although they require to be supplied with thiamin and other vitamins, which are normally supplied by the shoot in intact plants. Thus, different biosynthetic pathways are active in roots and shoots and it is clear that these pathways are strongly "locked".

It is commonly assumed that the cells in the shoot and root meristems are intrinsically uncommitted and that the different patterns of differentiation in the two organs lies in the structure and organization of the meristems themselves. However, an alternative hypothesis is that the meristematic cells of the shoot and root apices have become determined along "shoot" and "root" pathways of development respectively, i.e. that *intrinsic* differences in gene activity have been established, which are not caused primarily by the cellular environment in which they occur. This suggestion is in conflict with the current opinion that meristematic cells are completely uncommitted with respect to their future pattern of differentiation, but we have seen that in the phenomenon of phase-change, shoot apices can exist in two alternative but stable states (juvenile and adult). Since callus tissues derived from juvenile and adult shoots clearly exhibit intrinsic differences even when grown in isolation in aseptic culture (p. 321), it is possible that the differences in organization

and structure between juvenile and adult apices arises from the intrinsic, "determined" differences in the potentialities of their constituent cells.

On this basis it is logical to suggest that, by analogy, the characteristic structures and properties of root and shoot apices may arise from intrinsic differences in the developmental potentialities (though not in their *genetic* potentialities) of their constituent cells, rather than that the differences in the patterns of differentiation in root and shoot apices is the *result* of the organization and biochemical activities of the apices. Thus, the question at issue is whether the "locking" of meristematic cells into root or shoot pathways is a function of the multicellular structure and organization of the meristems, or whether it is an intrinsic function of the individual cells and is akin to other examples of determination. It is impossible on the present evidence to decide which of these alternative hypotheses is valid.

The establishment of root and shoot apices in the embryo is a major step in the overall differentiation of the plant body and further development involves differentiation at the organ, tissue and cell levels (p. 21). The question we now have to consider is whether unequal division and determination play essential roles in differentiation at these other levels.

We have seen that in lower plants having single apical cells, the proximal daughter cell is formed by unequal division of the apical cell and gives rise to mature tissue after a limited number of further divisions, and that it is possible to trace the cell lineages from the apical cell to the mature cells. It is difficult to identify the initial (promeristem) cells in roots of higher plants, but it is nevertheless possible, by studying the patterns of cell division, to trace the cell lineages back to the margins of the quiescent centre. Thus, the cell lineages which ultimately form specific tissues in the mature root (central cylinder, cortex, epidermis and root cap) can be seen to be established very close to the promeristem itself. However, we do not know whether the cells of any given lineage are already *determined* to give rise to certain tissues; that is to say, we do not know whether (1) the developmental potentialities of the early derivatives of the initial cells of the promeristem are already different and that these differences are transmitted through further divisions and so give rise to the various tissue zones of the mature root, or (2) the cell lineages are, at first, still capable of giving rise to any of the main zones of the root, and only later is the pattern of differentiation imposed on them by some unknown process.

The fact that in some roots cell lineages can apparently be traced back to specific layers of cells in the meristem region is consistent with the Histogen Theory put forward by Hanstein in 1868 according to which three meristematic zones or *histogens* can be recognized in both shoot and root apices, known as *plerome*, *periblem* and *dermatogen*, which were held to give rise to the vascular cylinder, cortex and epidermis, respectively. Thus, according to the Histogen Theory, initiating cells of the various tissue regions are separate, and destined to give rise to different regions of the mature root. It is possible, therefore, that the quiescent centre does, indeed, contain cells of different developmental potential.

Although the histogen concept is still a tenable hypothesis for root meristems, it does not appear to be applicable to shoot apices of higher plants for the following reasons. Firstly, the vascular tissue of leaf traces is formed from cortical cells and the pattern of development

of these traces appears to cut across any potential histogen boundaries that might be held to occur in the apical region of the shoot. Secondly, we have seen (p. 39) that the shoot apex has the properties of a self-organizing entity, with the capacity for self-regulation and regeneration. If this is so, then it would appear that the pattern of differentiation within the shoot apex is imposed on its constituent cells by the properties of the structure as a whole, whereas the hypothesis that differentiation involves an unequal division and the establishment of lineages of determined cells implies that the organization and properties of the apical region are determined by the intrinsic properties of its constituent cells.

Although it has been argued above that the difference between shoot and root apices may be dependent upon a degree of determination in the meristematic cells, it should be noted that this is not incompatible with the view that, so far as the differentiation of organs and tissues within the shoot apical region is concerned, it is the properties of the apex as a whole which regulate the pattern of differentiation rather than unequal division and the establishment of histogens.

The rather complex interactions between what might be referred to as the "intrinsic" and "organizational" influences affecting differentiation in shoot and root apices may be summarized as follows:

- (1) It is possible that the differences in structure between root and shoot apices depends upon intrinsic (determined) differences between their meristematic cells.
- (2) Organogenesis and tissue differentiation in the shoot apex are regulated by the organization and properties of the apex as a whole.
- (3) It is not clear whether differentiation in the root apex is partly regulated by its organizational properties, but the patterns of cell lineage are not inconsistent with the possible existence of histogens established close to the quiescent centre.

HORMONES AND DIFFERENTIATION

So far we have discussed mainly the role of intrinsic cell factors in differentiation, but we now have to consider the second situation referred to earlier (p. 316), namely that where the pattern of differentiation is controlled by factors *external* to the cell, and especially the role of hormones. By definition, the term hormone can be applied to a growth regulating substance only where it leaves the cell in which it is formed and affects another cell.

We have already met a number of instances where a hormone, or a combination of hormones, brings about a qualitative change which can be regarded as an aspect of differentiation, including:

- (1) the induction of buds and roots in callus tissue by the combined action of IAA and a cytokinin (p. 148);
- (2) the induction of vascular strands in callus tissue and in cortical and pith tissue in stems by IAA (p. 118);

- (3) the interaction of IAA and GA₃ in the differentiation of vascular tissue (p. 121);
- (4) The initiation of adventitious roots in stem tissue in response to IAA (p. 135).
- (5) the induction of flowering and control of sex expression by gibberellins, auxins and ethylene (p. 242).

Although these appear to be valid instances in which hormones appear to stimulate differentiation, most of the effects of hormones appear not to control selective gene expression directly, but rather to stimulate growth and differentiation in cells and tissues which are already "programmed" to differentiate in a specific manner. We have seen that each type of hormone has a wide range of effects and that the same hormone may have quite different effects in different tissues. For example, IAA may stimulate cell vacuolation in young fruits and in developing internodes, but it stimulates cell division in the cambium and it is necessary for the formation of the secondary wall in xylem differentiation. Again, GA₃ will stimulate internode elongation in many plants, and the synthesis of α -amylase in barley aleurone. Thus, *in plants the specificity of hormone action resides in the "target" tissue itself* and the hormone does not seem to determine the pattern of selective gene expression.

It also has to be borne in mind that only five main types of endogenous hormone have so far been identified, whereas during the life cycle of the plant differentiation must involve the regulation of large numbers of genes in the right cells in the right sequence, and it seems unlikely that the same small number of hormones could regulate the expression of so many genes. However, it is possible that the establishment of the major developmental pathways involves the regulation of certain "master genes" controlling the activities of a large number of subordinate genes, which become activated during the subsequent stages of differentiation. Indeed, it is a striking feature of certain aspects of differentiation that it appears to involve the co-ordinated expression of blocks of genes, as in the development of a leaf or a flower. The number of major steps in the development of a higher plant involving master genes is probably quite small and it is possible that interaction between the known hormones may play a controlling role at some of these steps.

For the reasons already given, it seems unlikely that the known hormones act as specific "effector" substances in gene expression and possibly the finer control of gene activity involves substances with a higher specificity of action, such as protein or RNA molecules. Indeed, there is evidence that hormones bind with specific proteins and, if this is the case, then the specificity of hormone action in any given type of target cell may depend upon the receptor protein rather than on the hormone molecule. It is possible, therefore, that determination involves the production of specific receptor proteins and that differentiation will subsequently follow a predetermined pattern when the hormones become available to bind with the proteins.

Animal embryologists apply the term *competence* to the capacity of cells to respond to a developmental signal in a predetermined manner. Although competence in plants is less well recognized, there seems little doubt that an analogous phenomenon occurs in plants. A particularly clear example is provided by the effects of hormones on the cambium. As

we have seen (p. 121), both IAA and GA_3 stimulate cambial division and IAA is necessary for the differentiation of the cambial derivatives into xylem elements, while phloem development is promoted by high GA_3 levels. However, the normal pattern of differentiation of the cambial derivatives into xylem on the inner side and phloem on the outer side is apparently not controlled by the hormones but reflects a difference in competence of the derivative cells on the two sides, since if a segment of the cambial region is excised from a *Ricinus* hypocotyl and replaced in the reverse position, the xylem is now formed on the outside and phloem on the inside (p. 120). Another good example of the phenomenon of competence is seen in the responses of specific cells on the flanks of the apical meristem of *Lolium temulentum* to a floral stimulus (Fig. 10.6). We have no idea how the competence of specific groups of such target cells is determined, and this remains yet another of the unsolved problems of development.

SHORT-RANGE CELL INTERACTIONS IN PLANTS

Cell interactions mediated via hormones may occur over short or very long distances, but we now have to consider short-range, cell-to-cell interactions in plants. In animals, short-range cell interactions are well established. Firstly, since animal cells are motile and different cell types may become intermingled it is necessary that they should have the capacity for *cell recognition*, to enable them to distinguish between cells of the same and of a different type from themselves. This function appears to be fulfilled by the occurrence of recognition substances, especially proteins, in the cell surfaces. Secondly, in animal embryos certain tissue types have the capacity for inducing differentiation in neighbouring cells with which they are in contact and there is strong evidence that this effect involves diffusible substances capable of moving from the inducing to the induced cells. By contrast, somatic plant cells are not motile and the need for the capacity for cell recognition is not apparent, but sexual reproduction in plants, on the other hand, does involve motile cells and there is now increasing evidence for cell recognition mechanisms in pollen/stigma interactions in higher plants.

The discovery of cell-recognition processes in plants arose from studies on incompatibility in flowering plants. Many plant species have genetically determined incompatibility systems which prevent self-fertilization and hence promote outbreeding. In *gametophytic* self-incompatibility there are one or more gene loci each with multiple alleles (S_1, S_2, S_3 , etc.) and the S allele borne by each pollen grain is expressed phenotypically when it alights on the diploid stigma, rejection occurring when the same allele occurs in both pollen and stigma (Fig. 13.10). In this type the pollen germinates and the tube penetrates the stigma, but growth is arrested either in the stigma or in the transmitting tissue of the style before fertilization can occur. In *sporophytic* self-incompatibility the behaviour of the pollen grains is determined by the genotype of the parent plant, and dominance may be demonstrated (Fig. 13.10). In this system, the "self" pollen grain starts to germinate

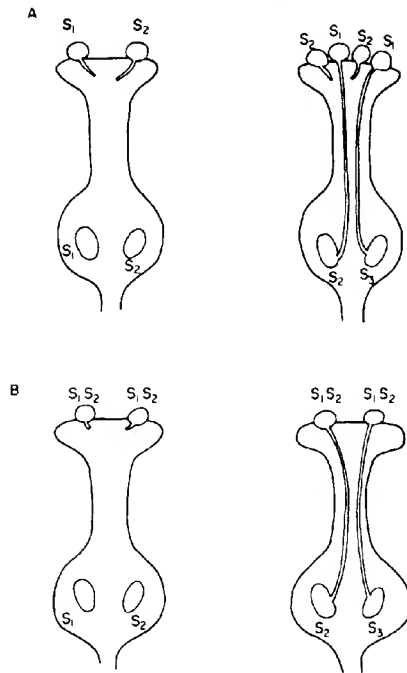


FIG. 13.10. Gametophytic (A) and sporophytic (B) self-incompatibility. In the gametophytic system, the S allele expresses its own phenotype during pollination on the diploid stigma, whereas in the sporophytic system, the phenotype of all pollen grains is determined by the genotype of the parent (S_1S_2). In both systems illustrated, the left-hand pollinations are self-incompatible and the right-hand ones compatible for (a) S_1 pollen grains of the gametophytic system, and (b) for all pollen grains of the sporophytic system, assuming that S_1 is dominant.

and the tube may even penetrate the stigma surface, but growth is arrested thereafter, so fertilization is again prevented.

The walls of pollen grains consist of two layers, an outer *exine* and an inner *intine*. In many species the exine is complex and includes cavities containing proteins, and the intine also contains proteins. The exine proteins are derived from the tapetum of the anther and so are sporophytic in origin, whereas the intine proteins are formed later in the pollen grain itself. When a pollen grain alights on a moist stigma surface, the exine proteins are released and diffuse into the stigma surface within a few minutes. It can be shown that the pollen exine proteins bind with proteins in the stigma surface. In the families Cruciferae and Compositae, with the sporophytic system, the pollen grain swells and germinates, but

in incompatible matings growth of the tube quickly stops, and the tip becomes occluded with callose (a β -1, 3-linked glucan). A counterpart reaction is induced in the contiguous stigma cells, which also produce callose deposits on the inner surface of the wall at the point of contact with the incompatible grain. The callose depositions are not formed in compatible matings, and the tube continues growth to effect fertilization. Even non-viable pollen can produce the callose responses, but washed pollen does not do so. Even extracts of pollen will produce the callose response and such extracts have been shown to contain glycoproteins. Thus, the reaction at the stigma surface can discriminate between wall proteins of different origin.

In plants showing the gametophytic system of self-incompatibility the response depends on interactions in the stylar canal or transmitting tissue, or in some cases, such as the Gramineae, in the stigma. Where the response is early, the intine-born proteins of the pollen grain may be involved, but where it is delayed the synthesis of the incompatibility factors probably takes place during the later growth of the tube. The interaction is with the interstitial material of the transmitting tract, which is rich in glycoproteins. The arrest of the tube in the Gramineae is accompanied by the deposition of callose, but there is no counterpart reaction in the stigma. In species with delayed tube inhibition, such as in the Solanaceae, the tube tip may become occluded, or it may burst with the release of particles containing callose precursors.

Little is known as to the nature of events occurring between the recognition reactions involving binding between pollen and stigma/style proteins and the subsequent rejection responses including the inhibition of tube growth and the deposition of callose. In the sporophytic system as exemplified by the Cruciferae, however, it seems likely that a diffusible substance is produced at the binding site on the stigma surface which stimulates the synthesis of callose, since this compound would not otherwise be formed in the stigma papillae. The response is in some respects similar to that induced in plant tissues by invading pathogens.

Although much remains to be elucidated regarding the nature of the pollen/stigma interactions it seems clear that the recognition of "self" from "not self" pollen is achieved through the initial binding reactions between pollen and stigma proteins.

It is not yet possible to say whether analogous recognition reactions occur between somatic cells of the plant body, but it may be significant that grafts between certain genotypes are successful and others unsuccessful. Little is known as to the molecular basis of acceptance or rejection between grafted tissues in plants, although a great deal is known regarding the corresponding phenomena in animals. It is possible, however, that cell recognition responses, analogous to those in pollen/stigma interactions, occur between plant somatic cells.

There is increasing evidence that glycoproteins play an essential role in cell-recognition phenomena in both plants and animals. Such glycoproteins are of widespread occurrence in plants, especially in seeds, but their function has hitherto remained unknown. From the evidence presented above, however, it seems very probable that the glycoproteins released from the exine of pollen grains function as recognition substances.

POLARITY AND PLANT FORM

It is self-evident that plant species show characteristic form and one aspect of this form is that there is typically a well-developed longitudinal axis, bearing lateral organs such as leaves and flowers. Differences occur along the axis, so that the two ends are not the same—for example, the plant axis is usually differentiated into a shoot end and a root end. In this respect the axis is said to show *polarity*, which has been defined as “any situation where two ends or surfaces in a living system are different”. Polarity of the axis of plants is most readily seen from morphological differences, but we have seen that it is also manifested in several physiological properties, as in the basipetal “polar” transport of auxin in stems, and in the regeneration of buds from the upper end and roots from the lower end of root segments (Fig. 13.13).

Although axial polarity is one of the most striking features of the plant body, it should not be forgotten that there are other forms of polarity. For example, the *dorsoventrality* of leaves involves differences between the upper and lower sides and may be regarded as a form of polarity. There may also be radial polarity in spherical bodies, such as cells of *Chlorella* or apple fruits, where there is a degree of radial symmetry, but there are differences between the inner and outer layers with respect to both chemical constituents and structure. In the following discussion we shall be concerned almost entirely with axial polarity and the term “polarity” will be taken to refer to this type, unless it is otherwise stated.

We have seen that polarization of the plant body into shoot and root in higher plants is initiated by the first unequal division occurring in the zygote. These unequal divisions are preceded by polarization of cytoplasmic components within the mother cell, and since the spindle of the first mitosis is orientated along the gradient of cytoplasmic components the first cross wall will result in an unequal distribution of these components between the two daughter cells. Thus, the polarity of the plant body throughout its development can be traced back to the initial polarization of the cytoplasmic components in the zygote. However, in the brown seaweed, *Fucus*, polarity is not already predetermined in the zygote, which thus provides favourable experimental material, since it is a free living cell in which the plane of the first division can be influenced by various treatments.

The eggs of *Fucus* are released into the surrounding seawater and after fertilization they settle in a solid stratum, where the development of a localized protuberance is the first sign of a rhizoid, by which the young plant becomes attached to a rock. The position of the rhizoid is determined by the orientation of the spindle in the first mitosis, which in turn determines the plane of the first dividing wall. The orientation of the spindle can be influenced by gradients in various external factors, such as light, heat, pH and osmotic activity. If the eggs are illuminated unilaterally, the shaded side (remote from an incident light) begins to form a protuberance at about 14 hours after fertilization and mitosis takes place so that the axis of the spindle is parallel in the direction of the incident light and a cell wall is formed at right angles to this, cutting off a larger cell which gives rise to a future thallus and a smaller cell which forms the rhizoid (Fig. 13.11). External factors can influence the axis of polarity up to a few hours before the appearance of the rhizoid, after which the position of emergence of the future rhizoid, appears to be irreversibly fixed.

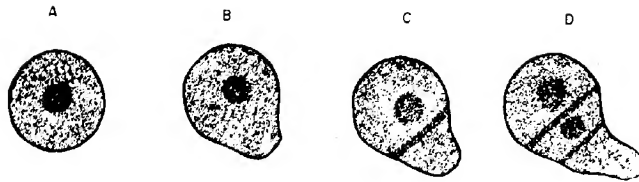


FIG. 13.11. Successive stages in the early development of the embryo of *Fucus*. In C, a wall has been formed giving rise to the rhizoid and thallus cells. In D, further division has occurred.

Before there are any visible signs of emergence of the rhizoid, the nuclear surface becomes highly polarized with finger-like projections (representing membranous extensions of the nuclear envelope) radiating towards the site of rhizoid formation. Mitochondria, ribosomes and fibrillar vesicles are also concentrated in this region of the rhizoidal half of the zygote, which is thus more densely cytoplasmic. At the same time changes occur in the cell wall at the future rhizoid side, with the deposition of the polysaccharide, fucoidin. As a result of these submicroscopic differences between the two halves of the cell, the two daughter nuclei come to lie in different cytoplasmic environments.

These submicroscopic changes are associated with changes in the electrical properties of the eggs. It has been shown that the future rhizoidal side of the cell becomes electronegative with respect to other parts, as a result of which a current is driven through the cell, with evidence that Na and Ca ions enter at the rhizoidal end and chloride ions at the thallus end. Moreover, the axis of polarity of an unpolarized egg can be fixed by applying a voltage across it. It has been suggested that the current helps to bring about differentiation as well as axis determination by an electrophoretic effect whereby negatively charged macromolecules are accumulated at certain points.

Light has a similar effect in determining the plane of the first cell division in the spores of the horsetail, *Equisetum*, and of other lower plants, the first cell wall being at right angles to the incident light (Fig. 13.12).

Once polarity has been induced it becomes extremely difficult or impossible to reverse it. This fact is well illustrated in regeneration experiments. Thus, if we take stem pieces of a plant such as willow and suspend them in a moist atmosphere, they will develop adventitious roots towards the morphologically lower end and the buds will tend to grow out most strongly at the upper end (Fig. 13.13). Similarly, if we take segments of the roots of chicory, dandelion or dock, and plant them in moist sand, roots will develop mainly from the morphologically lower (distal) end and buds from the upper (proximal) end (Fig. 13.13). Thus, although the willow cutting, and still less the root cutting, does not show any morphological differentiation between the upper and lower ends, nevertheless it is clear that these organs possess a marked physiological polarity which affects the pattern of

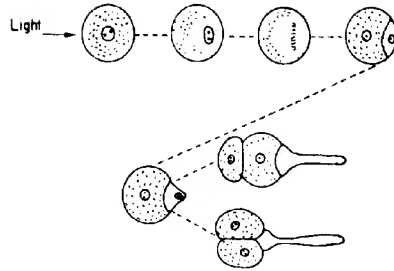


FIG. 13.12. Induction of polarity in a spore of *Equisetum* by unilateral illumination. (From D. von Wettstein, *Encycl. Plant Physiol.* **15**(1), 1965.)

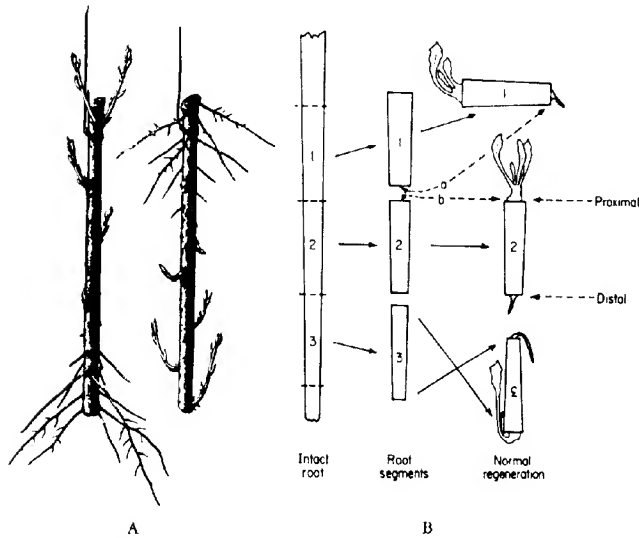


FIG. 13.13. A. Polarity of regeneration in willow stem. *Left*: stem cutting suspended in moist air in its normal orientation. *Right*: stem cutting similarly treated but in inverted position. Roots grew out at the morphologically lower end and shoot buds at the morphologically upper end, regardless of orientation. (Reprinted from *Pfeffer's Physiology of Plants*, 2nd ed., Clarendon Press, Oxford, 1903.)

B. Polarity of regeneration in root segments, such as those of *Taraxacum* and *Cichorium*. Shoot buds develop at the proximal end (i.e. the originally furthest from the root tip), and roots at the distal end, regardless of orientation. (Reprinted from H. E. Warmke and G. L. Warmke, *Amer. J. Bot.* **37**, 272, 1950.)

regeneration of roots and buds. This polarity is inherent in the tissues themselves and is not dependent on gravity, illumination or other external conditions, as is shown by the fact that willow cuttings suspended upside down in a moist atmosphere still regenerate roots predominantly at the original, morphologically lower end (Fig. 13.13), and similar effects are obtained with root segments of chicory which are planted so that the original upper end is now lowermost. Lower animals such as *Hydra* and planarians show analogous properties in the polarity of their regeneration patterns.

The physiological polarity of pieces of stem and root is a "built-in" property of the tissues. It might be thought that physiological polarity is due to gradients of metabolites or other substances along the stem or root, but this cannot be so, since polarity persists from one season to the next, through the dormant period, when metabolism is proceeding at a low rate, so that concentration gradients of metabolites are unlikely to persist. The basipetal transport of auxin (p. 100) and the occurrence of auxin gradients in the stem must therefore be seen as the *result* of the polarity of the tissues and not the cause of it.

We have seen that a piece of stem shows polarity of regeneration, with a tendency for buds to develop from the upper end and roots from the basal end. If such a stem is divided into two, each half behaves in a similar manner, and this process can be repeated even with very short pieces of stem. If it were possible to carry out this process to the limit, we might conclude that each individual cell exhibits polarity, and indeed there is evidence that this is the case. Thus, the filamentous green alga, *Cladophora*, shows polarity of the plant body, in that the basal end forms a rhizoid. If the cells of a filament of *Cladophora* are plasmolysed, so that the protoplasts are pulled away from the cell walls (which will break protoplasmic connections between cells), and are then deplasmolysed, each cell subsequently regenerates a new filament, developing a rhizoid at the basal end of the cell (Fig. 13.14). This experiment seems to provide clear evidence for polarity of individual cells of *Cladophora*. Comparable evidence for the cells of higher plants is more difficult to obtain, but much indirect evidence supports the hypothesis, such as the occurrence of unequal cell divisions.

These various observations suggest that (1) polarity of the tissues of a stem or root is remarkably stable and is not easily reversible, (2) polarity persists even throughout dormant periods, when metabolism is proceeding at a low rate, and (3) the polarity of a tissue apparently reflects the polarity of the individual cells. On these grounds, it has been suggested that there must be some permanent, structural basis for cell polarity. The observation that a piece of stem can be divided into several pieces, each of which shows the same polarity of regeneration, is very reminiscent of the fact that a bar magnet can be similarly divided and each piece becomes a small magnet. In a magnet each iron atom possesses magnetic polarity and the atoms become aligned during the process of magnetization, so that the "North" and "South" pole of each atom becomes oriented along the axis of the bar. By analogy, we might postulate that each cell of polarized organs, such as stems, contains polarized molecules which are oriented along the axis of each cell to form a "cytoskeleton", but attempts to demonstrate such oriented molecules in the cell have so far proved unsuccessful. It is interesting that the only linearly oriented structures that we do know of, the microtubules, lie at *right angles* to the axis of polarization in stems and roots (p. 9). It has been suggested



FIG. 13.14. Single cell from a filament of *Cladophora* regenerating a thallus from its apical end, and a rhizoid from the basal end. (Redrawn from A. T. Czaja, *Protoplasma*, **11**, 601, 1930.)

that polar transport of auxin may depend upon polarization of the plasma membranes at the end walls of the cell (p. 103).

Polarized Cell Division

The planes of cell division during the development of an organ play a very important role in determining its final form and shape. Indeed, we may say that without oriented cell divisions there could be no organized form within the plant body, as we see in cultures of callus tissue, where the plane of cell division is at random, and the resulting tissue forms a shapeless and structurally unorganized mass (p. 145). Thus, polarized cell divisions give the plant body its three-dimensional form. For example, in a developing internode the majority of cell divisions are oriented so that the mitotic spindle lies parallel to the axis of the internode. Consequently the internode grows mainly in length and relatively much less in diameter. For gourd fruits of different shape it has been shown that in long, elongated fruits divisions in which the mitotic spindle is oriented parallel to the long axis are much more frequent than divisions in which the spindle is in other planes, whereas in round fruits divisions in one particular plane do not predominate (Fig. 13.15). However, we cannot yet say what determines the planes of cell division along certain axes. It is clear that nuclear

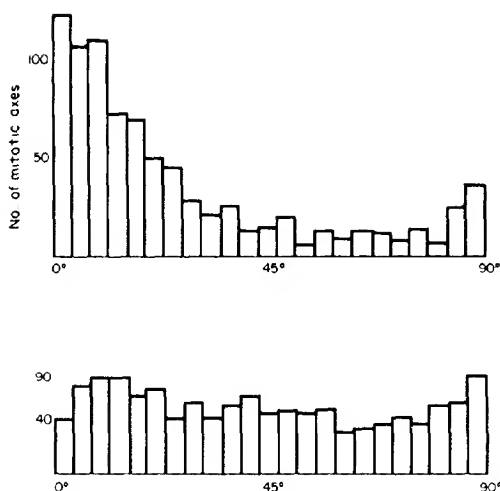


FIG. 13.15. Distribution of angles between mitotic spindles and longitudinal axis of the ovary in an elongate type of cucurbit fruit (*above*) and an isodiametric one (*below*). There is evidently a higher proportion of divisions nearly at right angles to the axis (spindles with low angles) in the former. In the latter, divisions are approximately equal at all angles. (From Sinnott, 1944.)

genes are involved, since many differences in form, including those affecting fruit shape in gourds, are inherited in a simple Mendelian manner.

It would appear that the axis of polarity of the whole organ also has a strong influence on the plane of cell division, by affecting the orientation of the mitotic spindle. How the orientation of the spindle is controlled is not known, but we have seen that it appears to be determined by the band of microtubules which appears in the cytoplasm before prophase, in the plane of the future cell plate (p. 9).

When we come to more complex forms, such as we see in a stamen or carpel, the problem of co-ordination of cell division and growth becomes much more difficult and at present we have no idea how this is achieved.

PATTERN IN DIFFERENTIATION

Patterns in the distribution of differentiated cells and tissues are very common and can take various forms. Thus, patterns may be seen in the arrangement of root hairs in regular longitudinal rows or in the regular markings seen in the petals of many flowers. Another example of pattern is uniform distribution of structures, such as leaf hairs, over a surface so

that they are separated from each other at approximately the same distance, forming a "mosaic". Other forms of pattern are manifold.

One factor which plays an important role in pattern formation is unequal division, as we have seen. Another phenomenon which seems to underlie pattern formation is the *mutual inhibition of like structures*. An example is seen in pattern formation of stomata in a developing leaf (Fig. 13.16). We find that the first series of stomata are uniformly distributed throughout the surface of the young leaf, but as this expands and the first formed stomata become separated by a greater distance, then new guard mother cells arise in the spaces between the original stomata. This pattern of behaviour suggests that developing structures of like nature, in this case guard mother cells, exert a mutually inhibitory influence on each other, so that when one structure occupies a given position, no similar structure can arise within a certain minimum distance from it. The cause of this inhibition between like structures is unknown but it has been suggested that it is due to competition between developing structures for specific substances required for their differentiation, so that if one structure arises at a given point it will tend to monopolize these substances within a certain radius, and so will prevent a second similar structure arising within its "inhibitory field".

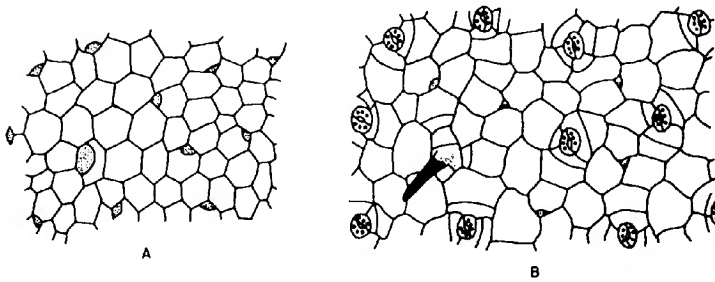


FIG. 13.16. Mutual inhibition between guard mother cells. Patterns of stomatal mother cells in a dicotyledonous leaf. A. The initial cells are shaded, but the large cell is a hair initial. B. Later stage in which, due to expansion of the leaf, these initial cells have moved further apart and developed into guard cells. The enlarged spaces between these allow the formation of new stomatal initials. (Redrawn from E. Bünning, *Survey of Biol. Progress*, Vol. 2, p. 105, Academic Press, New York, 1952.)

Very often isolated cells, such as guard mother cells, show cell division after the surrounding cells have ceased to divide and Bünning has called them *meristemoids*. He suggests that mutual inhibition is a property of such meristemoids and he has shown that a number of other examples of pattern can be interpreted in terms of this hypothesis. For example, he suggests that the formation of discrete strands rather than a continuous cylinder of procambial tissue in the stem is due to mutual inhibition of like structures, so that a strand can only develop at a certain minimum distance from a neighbouring strand. The origin of leaf primordia at the shoot apex may provide a further example of such inhibition (p. 31).

NON-GENIC FACTORS IN DEVELOPMENT

So far, we have regarded development as a process of "selective gene expression", involving the controlled activity of specific groups of genes, which in turn control the synthesis of enzyme and structural proteins characteristic of specialized cells. The information encoded in the DNA determines the amino acid sequence in the polypeptide chains of proteins, constituting their *primary* structure. Once the primary structure is established, the polypeptide chain can usually take on higher configurations without further directions from the genome. That is, genetic control of primary structure determines the secondary, tertiary and quaternary structure.

However, the cell is clearly not simply a random mixture of enzyme and other proteins, but has a highly ordered structure, in the form of organelles, membranes, etc., the structure of which appears not to be controlled by enzymic processes and hence not directly controlled by the information in the DNA. One way in which certain subcellular structures are formed is by the process of *self-assembly*, in which larger units, such as ribosomes, microtubules and membranes, are formed by the spontaneous assembly of smaller sub-units, frequently protein or nucleic acid macromolecules. This self-assembly is not directly enzyme controlled, but results directly from the physicochemical properties of the constituent sub-units.

Apart from the well-established process of self-assembly at the molecular level, there is other evidence of the importance of non-genic factors in development at the organelle and cell levels of organization. Thus, a considerable number of instances of "cytoplasmic inheritance" (in which certain characters of the offspring are transmitted through the maternal cytoplasm) are known for plants, and we now know that certain organelles, notably the plastids and mitochondria, have some degree of autonomy and indeed contain their own DNA.

Other non-genic factors probably include certain "physical" properties of cells and tissues. We have already seen the importance of external environmental factors, such as light and temperature, in plant development, but we must also consider the effects of other physical factors, such as surface tension and diffusion gradients of oxygen, arising and operating within the cells and tissues themselves. For example, attempts have been made to account for the position of cell walls in terms of physical laws. Earlier workers drew a parallel between the shapes and arrangements of cells in a tissue and those of groups of soap bubbles. Now, the shapes of soap bubbles can be interpreted in terms of the effects of surface tension, which tends to make them adopt forms with the least possible surface area (Fig. 13.17). Errera suggested that the shapes of cells can be similarly interpreted (Fig. 13.18), although it is questionable how far cell walls can be regarded as equivalent to the almost weightless films of soap bubbles. Moreover many cell types, such as cambium cells, do not appear to adopt a form with a minimal surface area. It is possible, however, that Errera's laws are applicable to isolated cells, or groups of cells, which will be free of the pressures and other influences from surrounding cells which occur in a tissue mass. Thus, the position of the walls formed during the early development of plant embryos may follow Errera's law and be determined by surface tension effects.

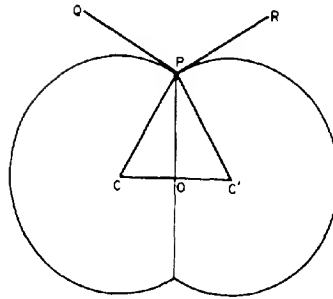


FIG. 13.17. Stable partition and walls of minimum surface assumed by two equal bubbles which are in contact. Angles OPQ and OPR are 120° . The distance between the centres equals the radii. (From D. A. W. Thompson, *Growth and Form*, Cambridge University Press, 1942.)

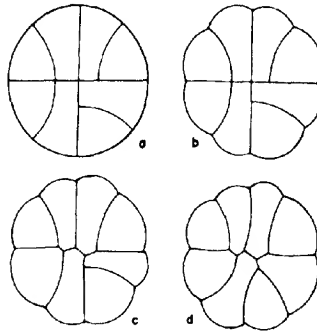


FIG. 13.18. Plate of eight cells (or bubbles) assuming a position of equilibrium where cell surfaces are of minimum area. (From D. A. W. Thompson, *Growth and Form*, Cambridge University Press, 1942.)

Although we cannot elaborate further on this theme here, it is clear that we should avoid the danger of assuming that all aspects of growth and development can be accounted for in terms of the information encoded in the DNA of the nucleus.

EPILOGUE

It will be apparent from the foregoing discussion of various general aspects of development that our knowledge is still very fragmentary and that although good progress is being made in certain areas there are others in which our understanding is almost totally deficient. For example, we have little precise information as to (1) how selective

gene expression is regulated in eukaryotes, (2) how cell heterogeneity is established in apical meristems, (3) how certain cells are "pre-programmed" to respond to a "signal" such as a growth substance, (4) what the "determination" of cells and tissues involves at the molecular level, (5) how regular spatial and temporal co-ordination of the processes involved in morphogenesis is achieved. Indeed, development presents some of the most important remaining unsolved problems in biology, the solution of which offers exciting challenges for the future.

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